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(54) Title: FUNCTIONAL SURROGATES OF ANALYTES OF INTEREST AND METHODS OF OBTAINING AND USING SAME

(57) Abstract

Functional surrogates are disclosed which serve as mimics of naturally occurring molecules, such as analytes of interest present in a given sample. In particular, functional surrogates (including epitopes and mimetopes) of macromolecular moieties, including large to medium-sized proteins, are described. The functional surrogates of the present invention are useful in a variety of applications, including diagnostic, prophylactic, and therapeutic applications. Indeed, where the detection of a macromolecular moiety is hampered by its size, a functional surrogate of the present invention, serving as the mimic of the macromolecular moiety, may be better suited for a given diagnostic assay. Methods of obtaining functional surrogates, various methods of their use, and compositions, including kits, are also described. Accordingly, certain binding peptides, including those of a well defined sequence, have been discovered, which can be used in a number of affinity assays, including those utilizing fluorescence polarization immunoassay (FPIA), enzyme multiplied immunoassay techniques (EMIT) or cloned enzyme donor immunoassays (CEDIA), to name a few.

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FUNCTIONAL SURROGATES OF ANALYTES OF INTEREST AND METHODS OF OBTAINING AND USING SAME

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1. Field of the Invention

The present invention relates to functional molecular surrogates of naturally occurring molecules, including a variety of analytes of interest, such as large to medium-sized proteins. The functional surrogates of the present invention can be used in a variety of applications, including diagnostic, prophylactic, and therapeutic applications. In particular, large macromolecular moieties whose detection may be impractical under certain assay conditions, such as the conditions of homogenous immunoassay techniques, are detected successfully with the aid of the functional surrogates of the present Methods of obtaining functional surrogates, various invention. methods of their use, and compositions, including kits, are also described. The invention also relates to certain constructs comprising DNA sequences encoding selected functional surrogates that exhibit the affinity and/or related characteristics required to mimic the function and/or behavior of the naturally occurring analyte molecules. transforming vectors including the constructs, in addition to bacteriophage and microorganisms harboring same.

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2. <u>Background of the Invention</u>

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The detection of various analytes that may be present in a given sample has always been of principal interest in science and medicine. The need for a method of determining the presence or absence of a given analyte of interest is particularly acute in a clinical

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setting, where assay conditions can be less than ideal, tensions especially high, and where speedy, reliable techniques may make the difference in the success or failure of the clinical treatment.

In most clinical settings, the assays for the detection of analytes of interest are indirect or of a heterogenous nature. Such heterogenous assays are time consuming and often require labeled antibodies for binding detection, not to mention solid carriers for use in separating bound from unbound antigenic species. Nonetheless, enzyme immunoassays (EIA) techniques are widely used for analyte detection because they are frequently the most effective of the available methods, or they may be the only method available for measuring the particular analyte of interest. See, for example, Porstmann, T. and Kiessig, S. T., in *J. Immunol. Meth.* (1992) 150:5-21, for a discussion of basic EIA techniques, including unlabeled (based on secondary immune reactions, such as precipitation and agglutination) and labeled (divided between so-called 'reagent-observed' and 'analyte-observed') methods. For the determination

of both haptens and high molecular weight substances, the authors favor the labeled method, which they characterize as using

monoclonal antibodies, as being of greater sensitivity, larger

Despite the great success enjoyed by enzyme immunoassays, artifacts and limitations persist. In particular, differences between the results of solution versus solid-phase techniques have been shown. See, e.g., Pesce, A. J. and Michael, J. B., in *J. Immunol. Meth.* (1992) 150:111-119. These differences are due to a number of factors such as surface phenomena, changes in molecular structure on binding to a surface, changes in the valence of antibodies and antigens, and steric

constraints. For other limitations of EIA, including sources of interference, the reader is referred to the article by Pesce and Michael, *supra*.

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2.1. "Homogenous" Affinity Assays

"Homogenous" enzyme immunoassays, those not requiring a surface bound component or a wash step, have been in use for a number of years since Rubenstein and co-workers described the inhibition of lysozyme activity on addition of morphine antibodies to a conjugate of morphine and lysozyme. Rubenstein, K. E., Schneider, R. S., and Ullman, E. F. *Biochem. Biophys. Res. Commun.* (1972) 47(4):846-851: U.S. Patent No. 4,190,496. When these workers discovered that the addition of free morphine reduced the inhibition of enzyme activity in proportion to the amount of free morphine added, the "homogenous" EIA technique was born.

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The extension of the technique was later shown for the detection of other haptens, including drugs and hormones, and to the use of other enzymes, such as malate dehydrogenase, glucose-6-phosphate dehydrogenase (G6DPH), amylase, and beta-galactosidase. Gibbons, I. et al. Anal. Biochem. (1980) 102:167-170. The extension of the technique to macromolecular antigens proved more difficult, however, and such assays were adversely affected by serum. Moreover, the intimate interaction between enzyme and bound antibody, which is responsible for the conformational effects that give rise to the inhibition of enzyme activity, is less intimate and in fact attenuated when the enzyme is bound to a large protein antigen. Indeed, binding to the large protein antigen may sterically inhibit the enzyme to begin with and prevent the enzyme from interacting with

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professed. See, Gibbons, I. et al., supra. Still others have tried to improve the sensitivity of homogenous EIA for the detection of macromolecular antigens by the use of modified labels, such as fluorogenic substrates for the enzyme. Armenta, R. et al., in Anal. Biochem. (1985) 146:211-219, describe an assay for serum ferritin using a beta-galactosidase-ferritin conjugate and dextran-linked beta-galactosyl-umbelliferone as enzyme substrate. A 1000-fold increase in sensitivity in going from a chromogenic substrate to a fluorogenic substrate was asserted. However, serum interference remained problematic due to the presence of antibody against beta-galactosidase in the patient samples.

2.2. Automated Assays

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In recent years the major trend in the field has been toward non-isotopic assays capable of being automated. See, for example, Gosling, J.P., in Clin. Chem. (1990) 36(8):1408-1427. That is, immunoassays can be run manually by technicians performing the reagent addition steps - "manual assays", or on automated instruments - "automated assays". Automated assays can be run on either dedicated immunoassay instruments or on existing clinical chemistry analyzers. Dedicated Immunoassay instruments are usually differentiated by the detection mode used to monitor the assay (e.g., chemiluminescence, fluorescence, particle counting) and the method used, as in the case for heterogeneous assay systems, to separate free and antibody bound labeled ligand. Additionally a dedicated instrument is limited in that it will only run assays formulated

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specifically for that technology and particular detection system.

Conversely, homogenous immunoassays, without need for a separation or wash step, are particularly well suited for running in a conventional automated clinical chemistry analyzer. See, Khanna, in *Principles and Practice of Immunoassay*, C.P. Price & D.J. Newman (Eds.), Stockton Press, New York (1991) pp. 326-364.

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The Enzyme Multiplied Immunoassay Technique (EMIT), popularized by Rubenstein et al., supra, can be run automatically on such clinical chemistry analyzers. As noted above, EMIT is a competitive homogeneous EIA in which an analyte is labeled with enzyme (most commonly a hapten conjugated to G6PDH). Binding of antibody to hapten G6PDH results in a decrease of G6PDH activity. A competition is set up between labeled and unlabeled hapten for a limited number of antibody binding sites. Increased amounts of hapten in the sample lead to less antibody available to bind to the labeled hapten, hence increased G6PDH activity is the result of increased concentration of hapten in the sample. The assay reagents readily lend themselves to being run on automated clinical chemistry analyzers and require only rate measurements at 340 nm for monitoring. Enzyme activity is monitored by measuring the rate of NADH formation at 340 nm; i.e., the assay only requires a regular UV detection system for measurement.

There has been no commercialized application of EMIT for the measurement of large analytes (e.g., proteins and other macromolecular moieties) because of the lack of a suitable G6PDH-macromolecule conjugate whose activity can be inhibited; that is, the conjugation of G6PDH to a large molecule will inherently render G6PDH inactive.

As stated previously, in an EMIT assay, binding of exogenous

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antibody to enzyme-labeled antigen results in a change (a decrease) in observed enzymatic activity. Labeled and unlabeled antigen compete for a limited number of antibody binding sites. Hence, the concentration of antigen in the sample is directly proportional to the concentration of free labeled antigen. Accordingly, the greater the concentration of antigen in the sample, the greater the observed enzyme activity.

The only commercialized EMIT assays have been for the measurement of haptenic molecules, such as drugs of abuse or therapeutic drugs. Efforts, such as those by Gibbons, I., et al., supra, and Armenta, R., et al., supra, to extend the assay to macromolecular antigens have met with limited success. Both efforts require reagent incubation times in the order of hours, suffer from serum interferences and, in the case of the ferritin assay, require a detector for measuring fluorescence. In other words these assays were substantially inferior in ease of use and performance. It follows that in terms of ease of use, the EMIT procedure would only be practical for assays of haptens - small molecules such as drugs, as larger molecules, such as polypeptides or proteins, would inherently inactivate the enzyme activity on conjugation to the G6PDH.

In addition, an attempt to generalize this technology to other proteins of commercial importance is limited by the fact that in a competitive assay format, substantial amounts of highly purified analyte are required for conjugation to the enzyme. For many proteins, this requirement is prohibitive.

2.3. Other "Homogeneous" Affinity Assays

Other homogeneous enzyme immunoassays have been

described. For example, Jenkins, S. H., in *J. Immunol. Meth.* (1992) 150:91-97, discusses in addition to EMIT, substrate-labeled fluorescence immunoassay (SLFIA), prosthetic group-labeled immunoassay (PGLIA) or apoenzyme reactivation immunoassay (ARIA), cofactor-labeled immunoassay, inhibitor-labeled immunoassay, and cloned enzyme donor immunoassay (CEDIA). All these techniques are susceptible to interferences present in the sample, however, as there is no wash step. The search for ways to measure large analytes continues.

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In the FETI technique, fluorescence excitation transfer immunoassay, the assay can be done in several configurations. The general principle is that two members of a binding pair are labeled. one with a fluorescein analog, the other with a rhodamine analog. The mixture is excited at the fluorescein absorption wavelength. If the labeled constituents are bound to each other, an energy transfer can take place and the fluorescein emission quenches the rhodamine This phenomenon permits an index of binding to be measured. In a relevant assay configuration, two distinct monoclonals are labeled. Energy transfer occurs only when the labels are brought into proximity by binding to the analyte. A fluorimetric analyzer has been designed and built to run a panel of FETI assays along with EMIT small molecule assays (reading NADH fluorescence). See, Ullman, E.F., Schwarzberg, M., Rubenstein, K.E., in J. Biol. Chem. (1976) 251(14):4172-8; Ullman, E.F., Khanna, P.Y., in Methods in Enzymology (1981) 74:28-60.

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Another major assay is enzyme channeling. The concept is to label each of two monoclonal (or polyclonal) antibodies with a different enzyme. The two enzyme labels are coupled in the sense

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that the product of one is a substrate for the second. In a specific instance, one antibody is labeled with glucose oxidase (GO). The second is labeled with horseradish peroxidase (HRP). The peroxide produced in the GO reaction is reduced by the HRP, resulting in oxidation of a leuco dye and production of a color. The coupled reactions go much faster when the two enzymes are held in proximity as when the antibodies to which they are attached form a complex with an antigen. The rate of color production is thus an index of analyte concentration. The principle could be demonstrated but has never worked well as the magnitude of the channeling effect is simply too small. See, Gibbons, I., et al., in Methods in Enzymology (1987) 136:93-103

An additional technique is called LOCI, which stands for luminescent oxygen channeling immunoassay. The method is based on the familiar concept of bringing together two species in order to initiate a measurable event. In this instance, the two species are beads coated with antibodies. The two are brought together in pairs by an antigen. In this respect, the technique is not dissimilar to latex agglutination. However, one set of particles is labeled with a photosensitizer dye and an "antenna" molecule. This arrangement is capable of exciting molecular oxygen, which diffuses the short distance to the second bead where it initiates a chemiluminescence process by exciting a special molecule coupled to the second bead. The result is that light is emitted when the beads are joined as a consequence of an antigen-antibody reaction. However, at low analyte concentrations, there is a nonspecific binding of the beads and, consequently, a nonspecific light emission. Furthermore, the technique requires special instrumentation and. although

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homogeneous, is not amenable to standard clinical analyzers. See, Ullman, E. F., et al., in *Proc. Natl. Acad. Sci. USA* (1994) 91(12):5426-30.

Yet another homogeneous assay is Microgenics' (Boehringer Mannheim) CEDIA. See, U. S. Patent Nos. 4,708.929, 5.120,653, 5.244,785, and 5.362.625. The method is based on the activity of the enzyme beta-galactosidase, which in this technique has been divided into two fragments. The acceptor fragment, EA, contains 97% of the enzyme's total mass. A smaller fragment, the donor (ED), is made by a recombinant DNA technique and contains on the order of about 80 amino acids. The ED can be engineered to contain lysine or cysteine groups at specified locations for linking.

In the CEDIA method, a hapten analog is attached to ED. Antibody binding the ED-hapten complex prevents its recombination with EA to form active enzyme. In this way, the enzyme activity is proportional to the amount of free hapten in a specimen. The sensitivity of CEDIA is perhaps one order of magnitude better than that for EMIT. For example, Microgenics has published on a vitamin B12 assay on the Cobas Mira with a sensitivity down to 100 pg/mL. See, Khanna, P.L. and Worthy, T.E., in *Diagnostics in the Year 2000*, Van Nostrand Rheinhold, Singh, P., Sharma, B.P., and Praveen, T. (Eds.) (1992) p. 2-38.

Efforts were made to apply CEDIA to high molecular weight analytes. An early effort involved binding TSH to the ED. An effect was demonstrated in this competitive format, but it was not large enough to be readily detected.

Another attempt came in the case of ferritin which has two kinds of repeating subunits. In this instance, an antibody to ferritin

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is conjugated to the ED. The conjugates bunch up around ferritin in the specimen and prevent the formation of the EA-ED complex. Unlabeled antibody is added along with the conjugate to further crowd the ED and provide increased steric hindrance to complexation. This assay has been introduced in a commercial format for the Hitachi 717 analyzer with a sensitivity of 50 ng/mL. The assay requires addition of a reagent containing Ab-ED and substrate to the specimen, followed by a 5-10 minute incubation step. A second reagent containing unlabeled antibody and EA is then added. After a 3-4 minute incubation step, the absorbance is read to give the final result. The methodology is claimed to be applicable to "analytes with multivalent antigenic determinants like CRP, hepatitis surface antigen."

Recent advancements have also been made in another "homogeneous" affinity assay technology known as fluorescence polarization immunoassay (FPIA). The technology is limited by the immeasurably small signal changes that occur when the analyte mass exceeds 20,000 daltons.

Wei, A-P. and Herron, J. N., in *Anal. Chem.* (1993) 65:3372-3377, describe the use of synthetic peptides as tracer antigens in FPIA techniques reportedly to detect high molecular weight antigens. In this work, a panel of 221 octapeptides of overlapping sequence designed to span all possible eight amino acid segments present in the two chains of human chorionic gonadotropin (237 amino acid residues between the two chains) was screened with a monoclonal anti-hCG antibody. A comparison of the binding affinity of a synthetic binding peptide, GSGSRLPGPSDTC (SEQ ID NO:75), derived from the structure of

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two binding peptides isolated from the panel. SRLPGPSD (SEQ ID NO:76) and RLPGPSDT (SEQ. ID NO: 77), showed that the synthetic binding peptide had a binding affinity constant (K_a) for the antibody of 1.6×10^{7} M⁻¹ versus a binding affinity constant of 4.8×10^{9} M⁻¹ observed for the naturally occurring hCG molecule. Hence, the synthetic peptide has a binding affinity that was more than two orders of magnitude lower than the naturally occurring molecule. Consequently, the synthetic peptide is ill-equipped to compete effectively with the natural hCG molecule for limited anti-hCG antibody: that is, the synthetic peptide is readily displaced by the natural hCG molecule in a competitive immunoassay format.

The Prosthetic-Group Labeled Enzyme Immunoassay (PGLEIA) is an assay in which apoglucose oxidase is inactive unless reconstituted by complexation with a ligand-labeled FAD (flavin adenine dinucleotide) analog. Antibody binding the FAD-Ligand conjugate prevents the recombination. Most of the work with this assay was done with haptens, and one study was reported in which an assay for IgG was demonstrated. See, Morris, D.L. et al., in *Anal. Chem.* (1981) 53:658-65.

In the FSIA (fluorogenic substrate-labeled immunoassay) method a ligand is covalently coupled to a fluorogenic molecule by an enzyme cleavable bond. When anti-ligand antibody is preoccupied with analyte in the specimen, the enzymatic cleavage reaction occurs and produces a fluorescent molecule. When no analyte is present, the antibody binds to the conjugate which sterically prevents the cleaving enzyme from acting. As a result, no fluorescent signal is produced. For example, the fluorogenic substrate can be a derivative of 4-methylumbelliferyl phosphate.

and the enzyme can be beta-galactosidase. Tests for IgG and IgM have been demonstrated. The IgG test has a sensitivity of 2 micrograms per ml. See, Worah, D., et al., in *Clin. Chem.* (1981) 27:673-677.

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2.4 Previous Efforts of Obtaining "Mimics"

Luzzago, A., et al., in *Gene* (1993) 128:51-57, describe the selection of nonapeptides from a random nonapeptide library which bind to the monoclonal antibody H107. Two consensus sequences were described, including YXXXXXW (SEQ ID NO:78) and GSXF (SEQ ID NO:79), in which position X is variable. The value of these sequences is unclear particularly because a competition experiment set up between a biotinylated synthetic peptide containing the first consensus sequence and recombinant human H-subunit ferritin provided anomalous results. In particular, the absorption reading attributable to mAb H107 bound to biotinylated synthetic peptide actually *increases* with the addition of competing analyte. A progressive *decrease* in the absorption signal would have been expected in a well behaved system. (See, Fig. 5 of Luzzago.) Significantly, no mention of affinity assays is made.

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Thus, previous efforts in the art to establish homogeneous assays have required reagent incubation times on the order of hours, have suffered from serum interferences and, in the case of the ferritin assay, required a specialized detector for measuring fluorescence. Most importantly, the existing technology allows the effective measurement of haptens, not macromolecular antigens. And where attempts have been made to measure macromolecular species, interferences and limitations persist which are not normally encountered with small hapten molecules. Also, no effective

substitutes have been discovered which faithfully reproduce the properties and characteristics of the macromolecular analytes to the point that existing techniques found effective for small molecules can be applied to the macromolecules of interest. Thus, no functional surrogates have been described, for instance, which can compete effectively with a given analyte, such as an antigen, for a limiting amount of affinity receptor, such as an antibody.

Accordingly, the present invention seeks to remedy the shortcomings in the state of the art of affinity assays, providing substances that can serve as functional surrogates of selected analytes of interest. Such substances are particularly useful for applications in the area of homogeneous immunoassays in which functional substitutes for "untractable" macromolecules are unavailable.

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3. <u>Summary of the Invention</u>

It is therefore an object of the present invention to provide functional surrogates of analytes of interest which for all intents and purposes serve as effective substitutes for the analytes of interest, particularly when the analytes are macromolecular moieties, the detection of which have to date proved unworkable within the framework of existing affinity assay technology, such as EMIT, CEDIA, fluorescence polarization methods, and the like. It is important to stress that in the methods of the present invention, actual knowledge of the molecular structure of the segment of the analyte of interest responsible for the affinity interaction with a receptor for the analyte is neither necessary nor essential. For a given analyte of interest, all that is needed is the availability of an

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affinity receptor having a selective affinity for the analyte (e.g., the availability of an antibody against an analyte of interest). Indeed, in some cases, as described further below, novel substances can be uncovered which show an affinity for an analyte of interest and which can be used as a receptor for the analyte. The formation of a complex between the novel substance and the analyte can then be detected.

A method is thus provided for determining the presence or absence of an analyte of interest in a sample by an affinity assay comprising: (a) providing (i) a labeled conjugate comprising at least one label attached to a functional surrogate capable of competing effectively with an analyte of interest for a limiting amount of an affinity receptor for the analyte, the labeled conjugate exhibiting an activity that is altered on interaction of the labeled conjugate with the affinity receptor and which activity can be measured and related to the presence or absence of the analyte in a given sample, and (ii) the affinity receptor: (b) combining the labeled conjugate and affinity receptor with a sample suspected of containing the analyte to provide a measure of the activity; (c) measuring the activity; and (d) relating the activity to the presence or absence of the analyte in the sample.

Various analytes of interest can be detected in this manner, from small haptens to large macromolecules, using available technology once the functional surrogate has been isolated and identified. Accordingly, specific embodiments of the present invention are directed to particular analytes of commercial importance, including various antigens and antibodies, and using various affinity assays well known to those of ordinary skill in the

art, including EMIT, CEDIA, and fluorescence polarization.

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Another object of the invention is to provide a homogeneous immunoassay kit comprising: (a) a labeled conjugate diposed in a first container means, the labeled conjugate comprising at least one label attached to a functional surrogate of an analyte of interest, the functional surrogate capable of competing effectively with the analyte for a limiting amount of an affinity receptor for the analyte, the labeled conjugate capable of exhibiting an activity that is altered on interaction of the labeled conjugate with the affinity receptor and which activity can be measured and related to the amount of the analyte present in a given sample; and (b) disposed in a second container means the affinity receptor and, optionally, any substance required for the labeled conjugate to exhibit the activity.

Consistent with the objective of the present invention, a functional surrogate of an analyte of interest is provided which comprises a peptide having an interactive group that allows the surrogate to compete effectively with the analyte for a limiting amount of an affinity receptor for the analyte. For the practice of the above-described methods, a further object of the present invention is to provide a labeled conjugate comprising at least one label attached to a functional surrogate of an analyte of interest, the functional surrogate capable of competing effectively with the analyte for a limiting amount of an affinity receptor for the analyte, the labeled conjugate capable of exhibiting an activity that is altered on interaction of the labeled conjugate with the affinity receptor and which activity can be measured and related to the amount of the analyte present in a given sample.

The invention also provides recombinant DNA constructs

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comprising a DNA sequence encoding a functional surrogate of an analyte of interest, the functional surrogate capable of competing effectively with the analyte for a limiting amount of an affinity receptor for the analyte. DNA constructs comprising a DNA sequence encoding a fusion protein of the functional surrogate of the invention are also contemplated, such as a fusion protein comprising the functional surrogate of the invention fused to the primary sequence of an enzyme label in the proximity of the label's active site. For example, the fused enzyme label can exhibit glucose-6-phosphate dehydrogenase activity.

Also provided for are transforming vector including the functional surrogate or fusion protein construct; a bacteriophage transformed by the vector encoding the functional surrogate and a microorganism transformed by the vector or infected by the bacteriophage.

Furthermore, it is an object of the present invention to provide a method of obtaining functional surrogate of an analyte of interest comprising: (a) selecting an affinity receptor exhibiting a selective affinity for an analyte of interest; (b) screening a random peptide library with the affinity receptor for a binding peptide; (c) isolating the binding peptide and identifying its primary structure. Moreover, functional surrogates of affinity receptors which exhibit a selective affinity for an analyte of interest can be obtained similarly by screening a random peptide library with the analyte.

In the method of the present invention, the identified peptide may further be prepared by known techniques, including solid phase synthesis and its capacity to compete with the analyte for a limiting amount of the affinity receptor confirmed or verified.

Alternatively, a synthetic binding peptide's capacity to selectively bind to an analyte can also be confirmed. Preferably, the binding peptide is isolated from a phage displayed random peptide library. The contents of such libraries can be designed and generated by known techniques.

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The above and other objects of the invention will become readily apparent to those of ordinary skill in the relevant art from the following detailed description and drawings, in which only the preferred embodiments of the invention are described and shown, simply by way of illustration of the best mode of carrying out the invention. As is readily recognized the invention is capable of modifications within the ordinary skill of the relevant art without departing from the spirit and scope of the invention.

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4. <u>Definitions</u>

To further assist those interested in practicing the invention, the following definitions are provided.

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Activity - Any detectable, measurable phenomenon attributable directly or indirectly to the action of a particular species, such as the enzymatic activity of an enzyme, the anticoagulant activity of heparin, the absorption spectrum of a product produced from a reaction mediated by the species in question, the emission spectrum of a fluorogenic compound, the color intensity produced in a chromogenic reaction mediated by the species in question, the current produced by an electrochemical transformation that can be related to the amount present of a species in question, the rate at which a given product is produced, photon emission, radioactivity, and the like. In the present

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invention, the activity will be attributable to action of the label or the labeled conjugate, infra.

Affinity Receptor - A molecule that exhibits a selective affinity for an analyte of interest as defined below. Hence, an affinity receptor of a given analyte will interact or bind selectively with that analyte in the presence of other potential binding partners. An example of a common affinity receptor is an antibody against a particular antigen or one of a pair of well known affinity couples, such as biotin-avidin or protamine-heparin.

Affinity receptors are preferably antibodies, both polyclonal and monoclonal, but can be any substance, protein, nucleic acid or saccharide that binds analyte selectively, preferably specifically. Antibodies are produced by introducing an immunogen into the bloodstream of a living animal. For a review of the production of antibody reagents, see, Hurn & Chantler in "Methods in Enzymology" Vol. 70, Part A, 1980 Academic Press, eds. Van Vunakis & Langone, pp. 104-142; Kohler & Milstein Nature (1975) 256: 495-497.

Analyte of Interest - Any substance whose detection is of interest to the practitioner. Such substances may constitute both small and large molecules, including but not limited to haptens, immunogens, drugs of abuse, therapeutic drugs, factors, cofactors, hormones, small and large antigens, various markers, immunoglobulins, specific antibodies, proteins, glycoproteins, polysaccharides, polynucleotides, lipopolysaccharides, other lipid-containing macromolecules, and the like. Ideally, affinity receptors of the analytes of interest are available. Most preferably, analytes of interest will be any molecule for which a peptide can act as a

functional surrogate in an assay using an affinity receptor, usually an antibody. Analytes can be any compound of interest ranging from small molecule drugs and haptens (MW 100 daltons) to large proteins (MWs up to 500,000 daltons) and infectious agents such as bacteria and viruses.

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Functional Surrogate - A substance that can serve as a mimic or substitute for a naturally occurring molecule, especially its functional aspects, such as the ability of that naturally occurring molecule to interact or bind selectively with an affinity receptor. Hence, a minimum requirement for a functional surrogate may be the capacity of the functional surrogate to compete with the naturally occurring molecule in question for a limited amount of affinity receptor. It is important to note that a functional surrogate may have a molecular structure (e.g., a primary sequence) that corresponds to a continuous or discontinuous epitope of a naturally occurring analyte. Alternatively, a functional surrogate may a molecular structure that differs substantially from that of the analyte or an immunoreactive group present in a segment of the analyte. Other characteristics or properties may be desirable in a given functional surrogate, including a much reduced molecular size relative to the naturally occurring molecule, a selective binding affinity (K_a) for an affinity receptor comparable to that of the naturally occurring molecule or conversely a dissociation constant (K_d) from an affinity receptor complex comparable to that of the naturally occurring molecule. The capacity of a functional surrogate to exhibit a competitive binding profile that comports to that obtained from the naturally occurring molecule may also be a desirable characteristic.

Label - Any type of marker that can be attached covalently or non-covalently to another moiety by which the presence of that moiety, such as a functional surrogate, can be detected or accounted for. The action of a label will give rise to a certain signal or activity, which can be measured. Labels can be radioisotopes, paramagnetic metals, fluorescent dyes, chemiluminescent markers, enzymes, colored or fluorescent particles (latex particles, glass beads, etc.), and the like.

Labeled Conjugate - A molecular entity that results from the molecular (e.g., protein fusion) or chemical (e.g., chemical linking) combination of a label and a functional surrogate. A labeled conjugate will have an activity associated with it, which activity becomes altered, i.e., either inhibited (decreased) or magnified (increased) on interaction of the labeled conjugate with another molecular entity, namely, an affinity receptor for a naturally occurring molecule. Like the unconjugated functional surrogate, the labeled conjugate should also be able to compete effectively with a naturally occurring molecule (typically, the analyte of interest) for a limiting amount of the affinity receptor.

Naturally Occurring- As used herein to describe a molecule, analyte or the like, "naturally occurring" can also encompass "unnatural" substances, such as those that are man-made, recombinant, non-endogeneous, non-indigenous or a pollutant, etc. The term "naturally occurring" is used merely to distinguish the analyte substance from the functional surrogate or labeled conjugate of the invention.

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5. <u>Brief Description of Drawings</u>

FIG. 1 presents the dose-response curve obtained from increasing concentrations of peptide SEQ ID NO:58, serving as a proposed functional surrogate of hepatitis B surface antigen (HBsAg). As described in the Examples section, the increasing amount of immobilized functional surrogate of HBsAg allowed greater proportions of a fixed amount of goat anti-HBsAg to be bound to a solid support, giving rise to an increase in the "activity" or optical density measurement taken after the addition of a second antibody conjugate, rabbit anti-goat IgG horseradish peroxidase, and appropriate HRP substrate.

FIG. 2 presents the results of competitive ELISA experiments that demonstrate the substantial similarity of the competitive binding profiles exhibited by immobilized functional surrogate versus that exhibited by immobilized naturally occurring antigen. These results also support the proposition that the functional surrogates of the present invention are capable of competing effectively with the naturally occurring analyte for a limiting amount of affinity receptor (e.g., antibody).

FIG. 3 presents the Ab dilution curve results for peptide bHEP11 (SEQ ID NO:74).

FIG. 4 presents the competitive ELISA results for peptide bHEP11 (SEQ ID NO:74).

FIG. 5 presents the Ab dilution curve results for peptide bHEP2-2 (SEQ ID NO:38).

FIG. 6 presents the competitive ELISA results for peptide bHEP2-2 (SEQ ID NO:38).

FIG. 7 illustrates a scheme for the generation of a random 8

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amino acid peptide library, R8C. Oligonucleotides were synthesized, converted into double-stranded DNA, cleaved with restriction enzymes, and cloned into the M13 vector, m663. The random peptide region is flanked by cysteine residues and is situated at the N-terminus of mature protein III.

FIG. 8 illustrates a scheme for the generation of another phage displayed random peptide library, R26, used in selected biopanning experiments.

FIG. 9 illustrates a scheme for the generation of the D38 phage displayed random peptide library.

FIG. 10 illustrates a scheme for the generation of the DC43 phage displayed random peptide library.

6. Detailed Description of the Invention

The invention relates to functional surrogates, most preferably binding peptides isolated from a random peptide library, useful as substitutes for a naturally occurring molecule that for one reason or another (such as undesirable size, unavailability, scarcity) cannot be used practically in a given application. Hence, having a functional surrogate in hand, certain methods can be performed, including, for example, homogenous enzyme immunoassays for large proteins. In addition, a host of other techniques previously only applicable to smaller molecules can now be carried out using the functional surrogate as a substitute for the analyte of interest. When the functional surrogate is able to compete effectively for an affinity receptor for the analyte, specific affinity binding interactions can be detected directly or indirectly, as the case may be.

By selecting a suitable affinity receptor with which to screen a random peptide library to isolate and identify binding peptides, functional surrogates of macromolecular analytes of interest can be obtained which mimic the binding properties, among other things, of the naturally occurring molecule. It is, thus, possible to design affinity assays, such as homogeneous EIAs to measure such macromolecules. After identification, the functional surrogates are synthesized using conventional techniques, including chemical synthesis, degradation of proteins, and in the case of peptides, optionally by recombinant techniques.

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to be measured.

In a specific embodiment of the present invention, peptide epitopes (i.e., peptides corresponding to a continuous epitope found in an antigen) are isolated and characterized from a random peptide library. In other cases, peptide mimetopes (i.e., those peptides having a molecular structure that differs from that found in a continuous epitope or those peptides having a molecular structure that is a composite of the structure of a discontinuous epitope) are isolated and characterized from a random peptide library. Hence, as used herein the term "mimetope" means peptides of a defined sequence which mimic the function of epitopes of macromolecules

In a preferred embodiment of the invention, peptides or mimetopes of a defined sequence may be used for formatting homogeneous enzyme immunoassays (EIAs) for measurement of analytes. The peptides represent specific epitopes on the analytes or mimetopes thereof which can be detected with specific affinity receptors, such as antibodies.

In general, the functional surrogates are then attached to a

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label suitable for the detection method of choice. For example, a peptide mimetope is conjugated to an enzyme exhibiting glucose-6phosphate dehydrogenase activity. The labeled peptide conjugate can then be exposed to and allowed to interact with polyclonal. monoclonal or bispecific antibodies or their fragments, such as Fab' or F(ab')2, which antibodies serve as receptors for the naturally occurring analyte. The complex formed from the interaction, typically a binding interaction, of labeled peptide with antibody results in the inhibition of enzyme activity. If the activity is monitored, then the observed activity can be related to the amount of analyte present in a given sample, especially when the observed activity is compared to that observed from at least one control (i.e., a sample with a known amount of analyte, such as below or above the detection limit to provide a negative or a positive control). Of course, either specific antibody against the analyte of interest or competing analyte in the sample can be detected in this manner. The reaction between labeled peptide conjugate and antibody, or fragment, is relatively selective, preferably specific, and takes place preferably, but not necessarily, at the antigen binding site on the antibody.

Peptides of 5-35, preferably less than 15, amino acid residues or more in length that include core sequences representing single epitopes, epitope composites or mimetopes thereof found in large molecules can be chemically synthesized. The desired peptide sequence is deduced from the nucleotide sequence of DNA inserts found in isolated phage clones from phage displayed random peptide libraries which bind to the target affinity receptor following the selection procedure, e.g., "biopanning" experiments. The target

affinity receptors are preferably polyclonal antibodies, most preferably specific monoclonal antibodies. In specific embodiments of the invention, individual functional surrogate peptides may contain from about 4 to about 100 amino acid residues. Still other peptides may have about 35 amino acids or less, such as 6-25 amino acids. The number of residues is somewhat variable because of the possible conformational requirements of the functional binding region of the surrogate and the need in some cases to have additional flanking sequences. Hence, preferred functional surrogates may have 8-14 amino acid residues, while others may have 8-20 amino acid residues (see, Tables 1 and 2). To reiterate, the use of the random peptide library means that the possible molecular structures of potential binding peptides are not limited to or dictated solely by the primary sequence of a proteinaceous analyte. Consequently, binding peptides can potentially be isolated corresponding to known epitopes, to previously unknown epitopes or to wholly unrelated but functionally equivalent structures of immunogenic analyte segments.

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TABLE I. SYNTHETIC FERRITIN PEPTIDES

(Note: all peptides bear free alpha amine and omega carboxyl, unless otherwise

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SEQ. ID. NO.	SEQUENCE	
1	SGGRALFQS	
2	*SGGRALFQS	
3	SGGRALFQS*	
4	*-eca-RGGRALFQS	
5	Ac-D-βala-βala-SGGRALFQS	
6	Ac-SGGRALFQS-βala-βala-D	
7	SGGRALFQS-βala-βala-D	
8	Ac-D-βala-Y-βala-SGGRALFQS	
9	Ac-SGGRALFQSD-βala-Y-βala	
10	Ac-SGGRALFQS-eCA-COOH	
11	Ac-RGGRALFQS-eCA-Y-eCA-D	
12	RGGRALFQSBBYBC	
13	SSINPTPSD	
14	*SSINPTPSD	
15	LRQPAVSGGR SLFQNLDPSR	
16	LROPAVSGGR SLFONLDSR	
17	RGGRALFQS-eca-KK	
18	KK-eca-RGGRALFQS	
19	*ESSALFQ	
20	*E-βala-SALFQS	
21	Ac-E-βaia-SALFQS	
22	SSLFQE	
23	•SSLFQE	
24	RAFFRD	
25	*RAFFRD	

26	KYGGMSLFQSQMTAGHHAGT
27	TAKEGSVGGASLFLELRAQC
28	ESSLFQ
29	ECSSLFQC
30	EGGASLF
31	ECGGASLFC

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TABLE 2. SYNTHETIC HEPATITIS PEPTIDES

(Note: all peptides bear free alpha amine and omega carboxyl groups, unless otherwise annotated. Also, all peptides with two cysteines are cylized as cystine.)

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SEQ. ID. NO.	SEQUENCE	
32	*CTGPRHLC	
33	*SDHPLYSR	
34 -	*LPGPPHLS*	
35	LPGPPHLS	
36	Ac-LPGPPHLS	
37	C-oK-LPGPPHLS C-oK-LPGPPHLS	
38	*C-oK-LPGPPHLS *C-oK-LPGPPHLS	
39	AcDC-eCA-LPGPPHLS AcDC-eCA-LPGPPHLS	
40	AcLPGPPHLS{E	
41	AcLPGPPHLS-Ok-{{-Ok	
42	*STTSIGPTK	
43	RCPSDGNCY	

	44	*RCPSDGNCY
	45	*PSDGN
	46	*RSPSDGNSY
	47	*CPSDGNC
5	48	*SPSDGN
	49	*CEEGAVLPKC
	50	*-eCA-CEEGAVLPKC
	51	CTKPSDGNYC
	52	*CTKPSDGNYC
10	53	Ac-oK-CTKPSDGNYC
	54	CTKPSSGNYC
	55	RCTKPSDGNYC
	56	*RCTKPSDGNYC
	57	eca-CTKPSDGNYC
15	58	*eca-CTKPSDGNYC
	59	CO-CH.————————————————————————————————————
		NH,
	60	CO-CH ₂ —— S
		NH-eCA-KTRPSDGNYC-CONH, biotin-NH.
	61	KCTKPSDGNCK
		,

62	*KCTKPSDGNCK
63	*KCTKPSDGNCKK
64	ECTKPSDGNCE
65	*ECTKPSDGNCE
66	@ECTKPSDGNCE
67	CTKPSDGNCK
68	*CTKPSDGNCK
69	*@CTRPSDGNYC
70	*CKPSDGNC
71	*CTKPSDGNC
72	*CPSDGNYC
73	*CKPSDGNYC
74	*@CTKPSDGNYC@Y

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For Tables 1 and 2, the "*" symbol refers to a biotin label: "@" or "eca" stands for epsilon amino caproic acid: "OK" refers to a "sideways" attached lysine, with the N-terminal peptide bond forming at the epsilon amino group; a "{" symbol refers to a branching lysine (MAP-"multiple antigenic peptide" technology discussed further below); "Ac" is an acetyl group: and intramolecular bonds, typically cystine groups, are indicated by the solid lines.

Accordingly, for example, SEQ, ID, No. 41 represents a tetrameric antigenic peptide, in which four copies of the peptide Ac-LPGPPHLS-OK (SEQ ID NO:369) are attached to the four animo groups of the poly-lysine core of a four-branch MAP represented by the symbol "{{" (MAP4). The structure of MAP4 attached at the C-terminal end to a "sideways" lysine (OK) is H₂NCH₂CH₂CH₂CH₂CH(NH₂)CONH

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The use of MAP technology to make high-density synthetic peptide systems has been described. See, for example, Tam. J.P., in *Proc. Natl. Acad. Sci. USA* (1988) 85:5409-5413; Tam. J.P. and Zavala, F. in *J. Immunolog. Meth.* (1989) 124:53-61; Briand, J.-P., et al., in *J. Immunolog. Meth.* (1992) 156:255-265; MAP technology has also been extended to solid phase synthesis on resins. See, for example, Applied Biosystems' *User Bulletin* (1992) No. 34

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Such small functional surrogates can be easily conjugated to labels, such as G6PDH by conventional techniques presently in use to couple haptens and drugs to comparable labels. Such coupling can be accomplished without appreciable loss of activity attributable to the label, and, hence, the same activity can be attributed to the labeled conjugate.

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The use of these peptide epitope/mimetope conjugates allows the construction of a variety of affinity assays, as mentioned earlier, including homogeneous EMIT-type, CEDIA, and "TDX" (fluorescence polarization) assays for the measurement of large polypeptides and proteins analytes.

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Examples of analytes which may be detected by the method of the invention include, but are not limited to ferritin, follicle stimulating hormone (FSH), thyroid-stimulating hormone (TSH), human growth hormone (hGH), immunoglobulin E (IgE), prolactin, parathyroid hormone (PTH), and human placental lactogen (HPL). In the area of fertility/pregnancy, human chorionic gonadotropin (hCG) and human luteinizing hormone (hLH) can be detected in addition to FSH. Infectious agents may also be assayed, including cytomegalovirus (CMV), chlamydia, streptomycin A, rubella, toxoplasma, herpes, and hepatitis. Also, the presence or absence of cardia markers, such as CK-MB, myoglobin, myosin light chain, and troponin, in addition to tumor markers, such as carcinoembryonic antigen (CEA), alpha-fetoprotein (AFP), PSA, and CA125 can be determined. The method of the invention also avails itself to rapid allergy screening.

The analytes of interest to this invention are broad and varied. They may be characterized by being monoepitopic or polyepitopic. The polyepitopic analytes will normally be poly (amino acids), i.e., polypeptides and proteins, polysaccharides, nucleic acids, and combinations thereof. Such combinations or assemblages include bacteria, viruses, chromosomes, genes, mitochondria, nuclei, cell membranes, and the like.

For the most part, the polyepitopic analytes employed in the subject invention will have a molecular weight of at least about 5,000 more usually at least about 10,000. In the poly(amino acid) category, the poly(amino acids) of interest will generally be from about 5,000 to 5,000,000 molecular weight, more usually from about 20,000 to 1,000,000 molecular weight; among the hormones

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of interest, the molecular weights will usually range from about 5,000 to 60,000 molecular weight.

The following are classes of proteins that are related by structure and are potential analytes of interest: protamines, histones, albumins, globulins, scleroproteins, phosphoproteins, mucoproteins, chromoproteins, kipoproteins, nucleoproteins, glycoproteins, unclassified proteins, e.g. somatotropin, prolactin, insulin, pepsin.

In addition, a number of proteins found in the human plasma are important clinically and include: Prealbumin, Albumin, α_1 -Lipoprotein, α_1 -Acid glycoprotein, α_1 -Antitrypsin, α_2 -Glycoprotein, Transcortin. Postalbumin, α_1 -glycoprotein, α_{1x} -Glycoprotein. Thyroxin-binding globulin, Inter-a-trypsin-inhibitor, Gc-globulin, Haptoglobin. Ceruloplasmin, Cholinesterase, α_2 -Lipoprotein(s), α_2 -Macroglobulin, α_2 -HS-glycoprotein, Zn- α_2 -glycoprotein, α_2 -Neuramino-glycoprotein, Erythropoietin, B-lipoprotein, Transferrin, Hemopexin, Fibrinogen, Plasminogen, B2-glycoprotein I, B2glycoprotein II, Immunoglobulin G. (IgG) or ,G-globulin, Mol. formula: $\gamma_2 \kappa_2$ or $\gamma_2 \lambda_2$, Immunoglobulin A (IgA), or A-globulin. Mol. formula: $(\alpha_2\kappa_2)^n$ or $\alpha_2\lambda_2)^n$, Immunoglobulin M, (IgM) or Mglobulin, Mol. formula: $\mu_2 \kappa_2$)⁵ or $(\mu_2 \lambda_2)^5$, Immunoglobulin D (IgD), or "D-Globulin ("D), Mol. formula: $(\delta_2 \kappa_2)$ or $(\delta_2 \lambda_2)$, Immunoglobulin E (IgE), or $_{1}$ E-Globulin (,E), Mol. formula: $(\epsilon_{2}\kappa_{2})$ or $(\epsilon_2\lambda_2)$, Free κ and γ light chains. Complement factors: C^1 (C^1q. C'1r, C'1s), C'2, C'3 ($\beta_1 A$, $\alpha_2 D$), C'4, C'5, C'6, C'7, C'8, C'9. Potential analytes of interest, such as:

BLOOD	CLOTTING FACTORS
International designation	Name

1	Fibrinogen
II	Prothrombia
Ha	Thrombin
III	Tissue thromboplastin
V and VI	Proaccelerin, accelerator globulin
VII	Proconvertin
VIII	Antihemophilic globulin (AHG)
IX	Christmas factor, plasma thromboplastin component (PTC)
X	Stuart-Prower factor, autoprothrombin III
XI	Plasma thromboplastin antecedent (PTA)
XII	Hagemann factor
XIII	Fibrin-stabilizing factor

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Important protein hormones include: Peptide and Protein Hormones such as Parathyroid hormone. (parahormone), Thyrocalcitonin, Insulin, Glucagon, Relaxin, Erythropoietin, Melanotropin, (melanocyte-stimulating hormone; intermedin), Somatotropin, (growth hormone), Corticotropin, (adrenocorticotropic hormone), Thyrotropin, Follicle-stimulating hormone, Luteinizing hormone, (interstitual cell-stimulating hormone), Leuteomammotropic hormone, (luteotropin, prolactin),

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Tissue Hormones, Secretin, Gastrin, Angiotensin I and II. Bradykinin, Human placental lactogen: and Peptide Hormones from the Neurohypophysis, such as Oxytocin, Vasopressin, Releasing

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factors (RF). CRF. LRF. TRF. Somatotropin-RF. GRF. FSH-RF,

Gonadotropin, (chorionic gonadotropin);

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PIF. MIF.

Other macromolecular analytes of interest are mucopolysaccharides and polysaccharides.

Illustrative antigenic polysaccharides derived from microorganisms are as follows:

Species of Microorganisms	Hemosensitin Found in
Streptococcus pyogenes	Polysaccharide
Diplococcus pneumoniae	Polysaccharide
Neisseria meningitidis	Polysaccharide
Neisseria gonorrhoeae	Polysaccharide
Corvnebacterium diphtheriae	Polysaccharide
Actinobacillus mallei; Actinobacillus whitemori	Crude extract
Francisella tularensis	Lipopolysaccharide Polysaccharide
Pasteurella pestis	
Pasteurella pestis	Polysaccharide
Pasteurella multocida	Capsular antigen
Brucella abortus	Crude extract
Heamophilus influenzae	Polysaccharide
Haemophilus pertussis	Crude
Treponema reiteri	Polysaccharide
Veillonella	Lipopolysaccharide
Erysipelothrix	Polysaccharide
Listeria monocytogenes	Polysaccharide
Chromobacterium	Lipopolysaccharide

Mycobacterium tuberculosis	Saline extract of 90% phenol extracted mycobacteria and polysaccharide fraction of cells and tuberculin	
Klebsiella aerogenes	Polysaccharide	
Klebsiella cloacae	Polysaccharide	
Salmonella typhosa	Lipopolysaccharide Polysaccharide	
Salmonella typhi-murium; Salmonella derby	Polysaccharide	
Salmonella pullorum		
Shigella dysenteriae	Polysaccharide	
Shigella flexneri		
Shigell sonnei	Crude. polysaccharide	
Rickettsiae	Crude extract	
Candida albicans	Polysaccharide	
Entamoeba histolytica	Crude extract	

The microorganisms which are assayed may be intact, lysed, ground or otherwise fragmented, and the resulting composition or portion, e.g., by extraction, assayed. Microorganisms of interest include: Corynebacteria (Cornebacterium diptheriae), Pneumococci (Diplococcus pneumoniae), Streptococci (Streptococcus pyogenes, Streptococcus salivarus), Staphylococci (Staphylococcus aureus, Staphylococcus albus) Neisseriae (Neisseria meningitiais, Neisseria gonorrheae) Enterobacteriaciae, (Escherichia coli, Aerobacter aerogenes, Klebsiella pneumoniae, Salmonella typhosa, Salmonella chloeracsuis, Salmonella typhimurium, Shigella dysenteriae, Shigella schmitzii, Shigella arabinotarda, Shigella fiexneri, Shigella

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boydii, Shigella Sonnei);

Other enteric bacilli. (Proteus vulgaris. Proteus mirabillis, Proteus morgani. Pseudomonas aeruginosa. Alcaligenes faecalis. Vibrio cholerae); Hemophilus-Bordetella group (Hemophilus influenzae. H. ducrevi, H. hemophilus. H. aegypticus, H. paraiufluenzae. Bordetella pertussis). Pasteurellae (Pasteurella pestis. Pasteurella tulareusis), Brucellae (Brucella melitensis Brucella abortus. Brucella suis), Aerobic Spore-forming Bacilli (Bacillus anthracis, Bacillus subtilis, Bacillus megaterium, Bacillus cereus), Anaerobic Spore-forming Bacilli (Clostridium botulinum, Clostridium tetani Clostridium perfringens. Clostridium novyi, Clostridium septicum. Clostridium histolyticum. Clostridium tertium, Clostridium bifermentans. Clostridium sporogenes);

Mycobacteria (Mycobacterium tuberculosis hominis, Mycobacterium bovis, Mycobacterium avium, Mycobacterium leprae, Mycobacterium paratuberculosis), Actinomycetes (fungus-like bacteria) (Actinomyces israelii, Actinomyces bovis, Actinomyces naeslundii, Nocardia asteroides, Nocardia brasilinesis);

The Spirochetes (Treponema pallidum, Spirillum minus, Treponema pertenue, Streptobacillus moniliformis, Treponema carateum. Borrelia recurrentis, Leptospira icterohemorrhagiae, Letospira canicola), Mycoplasmas (Mycoplasma pneumoniae, Other pathogens (Listeria monocytogenes, Erysipelothrix rhusiopathiae, Streptobacillus moniliformis, Donvania granulomatis, Bartonella bacilliformis), Rickettsiae (bacteria-like parasites) (Rickettsia prowazekii, Rickettsia mooseri, Rickettsia rickettsii, Rickettsia conori, Rickettsia australis, Rickettsia sibiricus, Rickettsia akari, Rickettsia tsutsugamushi, Rickettsia burnetii, Rickettsia quintana);

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Chlamydia (Chlamydia agents). Fungi, Cryptococcus neoformans. Blastomyces dermatidis, Histoplasma capsulatum. Coccidioides immitis. Paracoccidioides brasiliensis, Candida albicans, Aspergilus fumigatus, Mucor corymbifer (Absidia corymbifera), Rhizopus oryzae, Rhizopus arrhizus, Rhizopus nigricans, Sporotrichum schenkii, Fonsecaea pedrosoi, Fonsecaea compacia, Fonsecaea dermatitidis, Cladosporium carrionii, Phialophora verrucosa, Aspergillus nidulans, Madurella mycetomi, (Madurella grisca, Allescheria boyaii, Phialosphora jeanselmei, Microsporum gypseum, Trichophyton mentagrophytes, Keratinomyces ajelloi, Microsporum canis, Trichophyton rubrum. Microsporum andouini);

Viruses, Adenoviruses, Herpes viruses, (Herpes simplex, Varicella (Chicken pox), Herpes Zoster (Shingles), Virus B, Cytomegalovirus), Pox Viruses, (Variola (smallpox), Vaccinia, Poxvirus bovis, Paravaccinia, Molluscum contagiosum),

Picronaviruses (Poliovirus, Coxsackievirus, Echoviruses, Rhinoviruses):

Myxoviruses (Influenza (A. B and C), Parainfluenza (1-4), Mumps Virus, Newcastle Disease Virus, Measles Virus, Rinderpest Virus, Canine Distemper Virus, Respiratory Syncytial Virus, Rubella Virus), Arboviruses (Eastern Equine Eucephalitis Virus, Western Equine Eucephalitis Virus, Sindbis Virus, Chikungunya Virus, Semliki Forest Virus, Mayora Virus, St. Louis Encephalitis Virus, California Encephalitis Virus, Colorado Tick Fever Virus, Yellow Fever Virus, Dengue Virus), Reoviruses (Reovirus Types 1-3), Hepatitis (Hepatitis A Virus, Hepatitis B Virus, Gross Virus, Tumor Viruses (Rauscher Leukemia Virus, Gross Virus,

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Maloney Leukemia Virus).

The monoepitopic ligand analytes will generally be from about 100 to 2,000 molecular weight, more usually from 125 to 1,000 molecular weight. The analytes of interest include drugs, metabolites, pesticides, pollutants, and the like. Included among drugs of interest are the alkaloids. Among the alkaloids are morphine alkaloids, which includes morphine, codeine, heroin, dextromethorphan, their derivatives and metabolites; cocaine alkaloids, which includes cocaine and benzoyl ecgonine, their derivatives and metabolites; ergot alkaloids, which includes the diethylamide of lysergic acid; steroid alkaloids; iminazole alkaloids; quinoline alkaloids; which includes quinine and quinidine; diterpene alkaloids, their derivatives and metabolites.

The next group of drugs includes steroids, which includes estrogens, gestrogens, androgens, adrenocortical, bile acids, cardiotonic glycosides and aglycones, which includes digoxin and digoxigenin, saponins and sapogenins, their derivatives and metabolites. Also included are the steroid mimetic substances, such a diethyl stilbestrol.

The next group of drugs comprise cyclic lactams having from 5 to 6 membered rings, which include the barbiturates. diphenyl hydantoin, and their metabolites.

The next group of drugs is aminoaklyl benzenes, with alkyl of from 2 to 3 carbon atoms, which includes the amphetamines, catecholamines, which includes ephedrine, L-dopa, epinephrine, narceine, papaverine, their metabolites and derivates.

The next group of drugs is benzheterocyclics which include

oxazepam, chlorpromazine, tegretol, imipramine, their derivatives and metabolites, the heterocyclic rings being azepines, diazepines and phenothiazines.

The next group of drugs is purines, which includes theophylline, caffeine, their metabolites and derivatives.

The next group of drugs includes those derived from marijuana, which includes cannabinol and tetrahydrocannabinol.

The next group of drugs includes the vitamins such as A, B, C, D, E and K.

The next group of drugs is prostaglandins, which differ by the degree and sites of hydroxylation and unsaturation.

The next group of drugs is antibiotics, which include penicillin, chloromycetin, actinomycetin, tetracycline, terramycin, their metabolites and derivatives.

The next group of drugs is the nucleosides and nucleotides, which include ATP, NAD, FMN, adenosine, guanosine, thymidine, and cytidine with their appropriate sugar and phosphate substituents.

The next group of drugs is miscellaneous individual drugs which include methadone, meprobamate, serotonin, meperidien, amitriptyline, nortriptyline, lidocaine, procaineamide, acetylprocaineamide, propanoloi, griseofulvin, butryophenones, antihistamines, anticholinergic drugs, such as atropine, their metabolites and derivatives.

The next group of compounds is amino acids and small peptides which include thyroxin, triiodothyronine, oxytocin, ACTH, angiotensin, gentamycin, met- and leu-enkephalin their metabolites and derivatives.

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Metabolites related to diseased states include spermine, galactose, phenylpyruvic acid, and porphyrin type 1.

Among pesticides of interest are polyhalogenated biphenyls, phosphate esters, thiophosphates, carbamates, polyhalogenated sufenamides, their metabolites and derivatives.

For receptor analytes, the molecular weights will generally range from 10,000 to 2 x 10°, more usually from 10,000 to 10°. For immunoglobulins IgA, IgG, IgE and IgM, the molecular weights will generally vary from about 160,000 to about 10°.

Enzyme analytes will normally range from about 10.000 to 600,000 in molecular weight. Natural receptors vary widely, generally being at least about 25,000 molecular weight and may be $10^{\rm e}$ or higher molecular weight including such materials as avidin, thyroxine binding globulin, thyroxine binding prealbumin, transcortin, etc.

In addition, numerous hybridomas have been deposited and are available from the ATCC. Such hybridomas produce antibodies that can serve as affinity receptors for use in biopanning experiments to identify functional surrogates of specific antigens. See, for example, ATCC catalog of Cell Lines & Hybridomas, 7th Ed. (1992). See, for example, pp. 319-332 (secreted mAb).

Specific peptides of a defined sequence can be produced in vitro by synthesis or by chemical or enzymatic cleavage.

Alternatively, such peptide can be produced in vivo by a natural process. The peptides preferably have a MW of about 2000 or less, and must be capable of competing effectively in the presence of naturally occurring analyte for a limited amount of affinity receptor (e.g., antibody). The peptide should preferably exert only a

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minimal effect. if any, on the activity of a label when conjugated to that label. As stated elsewhere, labels can comprise fluorescent markers, enzymes, enzyme substrates, and the like. Examples of fluorescent markers include fluoresceins, rhodamines, cyanins, eosins, and the like. A preferred enzyme is glucose-6-phosphate dehydrogenase. Other suitable candidate enzymes are lysozyme or beta-galactosidase. A suitable candidate enzyme is one whose activity is little affected by conjugation to the functional surrogate, but is greatly affected by binding of affinity receptor to the labeled conjugate.

Table 3 lists some additional enzymes that may be suitable for use in the present invention, along with their enzyme substrates.

TABLE 3. LIST OF ENZYMES AND THEIR SUBSTRATES

5	I. <u>Hydrolases Carbohydrases</u>	
	Amylase, Lactase, Maltase, Sucrase, Emulsion	Carbohydrates. Starch. dextrin, etc. Lactose, Maltose, Sucrose B-Glucosides and derivatives
10	II. <u>Nucleases</u>	
	Polynucleotidase, Nucleotidase	Nucleic acid, Nucleotides
15	III. <u>Arginase</u>	
20	Arginase. Urease. Glutaminase, Transaminase	Amino compounds and amides, Arginine, Utea, Glutamine, Glutamic acid, etc.
	IV. <u>Purine Deaminases</u>	
	Adenase. Guanase	Adenine, Guanine
25	V. <u>Peptidases</u>	•
30	Aminopoluypeptidase, Carboxypeptidase, Dipeptidase, Prolinase VI. <u>Proteinases</u>	Polypeptides, Dipeptides, Proline peptides
35	Pepsin, Trypsin. Cathepsin, Rennin. Chymotrypsin, Papain, Ficin	Proteins, proteoses, Casein proteins, peptones
	VII. <u>Esterases</u>	
40	Lipase, Esterases. Phosphatases, Sulfatases, Cholinesterase	Fats, ethyl buryraate, etc., esters of phosphoric acid, esters of sulfuric acid, Acetylcholine

	VIII. Iron Enzymes	
5	Catalase. Cytochrome oxidase, Peroxidase	Hydrogen peroxide, reduced cytochrome C in the presence of oxygen, a large number of phenols, aromatic amines, etc., in the presence of H_2O_2
10	IX. <u>Copper Enzymes</u>	
	Tyrosinase (poly-phenoloxidase, mono-phenoloxidase, Absorbic acid oxidase	Various phenolic compounds, Ascorbic acid in the presence of oxygen
15	X. Enzymes Containing Co-Enzymes I and/or II	
20 25	Alcohol dehydrogenase. Malic dehydrogenase. Isocriterie dehydrogenase. Lactic dehydrogenase. B-Hydroxybutyric dehydrogenase, Glucose dehydrogenase, Glycerophosphate dehydrogenase, Aldehyde dehydrogenase	Ethyl alcohol and other alcohols, L() Malic acid, L-Isocritic acid, Lactic acid, L-\(\beta\)-Hydroxybutyric acid, D-Glucose, Robinson ester (hexose-6-phosphate) Glycerophosphate, Aldehydes
	XI. <u>Enzymes Which Reduce</u>	Cytochrome
30	Succinic dehydrogenase (as ordinarily prepared)	Succinic acid
	XII. Yellow Enzymes	
35	Warburg's old yellow enzyme Diaphorase, Haas enzyme, Xanthine oxidase, D-amino acid oxidase, L-Amino acid oxidases, TPN-Cytochrome	Reduced co-enzyme II. Hypoxanthine xanthine, aldehydes, reduced co-enzyme 1, etc., D-amino acids+ 0, L amino acids, reduced co-
40	C reductase, DPN- Cytochrome reductase	enzymes II and cytochrome C, reduced co-enzymes II and cytochrome C

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XIII. <u>Hvdrases</u>

5	Furnarnse, Aconitase Enolase	Fumaric acid + H ₂ 0 Citric acid, 2-Phosphoglyceric acid
	XIV. Mutases	() () () () () ()
10	Glyoxalase substituted glyoxals	Methyl glyoxal and other
	XV. <u>Desmolases</u>	
15	Zymohexase (aldolase), Carboxylase, ß-Keto carboxylases, Amino acid decarboxylases, Carbonic anhydrase	Fructose 1,6-diphosphate, pyruvic acid, ß-Keto acids L-Amino acids, Carbonic acid
20	XVI. Other Enzymes	
25	Phosphorylase, Phosphohexdo- isomerase, Hexokinase Phosphoglucomutase	Starch or glycogen and phosphate, Glycose-6-phosphate, Adenosine-triphosphate, Glucose-1-phosphate

As mentioned previously, specific binding peptides having a defined sequence can be obtained in a variety of ways, including isolation, subsequent to specific or non-specific enzyme digestion or chemical degradation of macromolecule; in vivo or in vitro production by transformed cells, tissue culture, and transgenic animals.

The specific binding peptide, such as any one of those listed in Tables 1 and 2, can then be conjugated by conventional methods to a label, preferably an enzyme. Conjugation methods such as those disclosed in U.S. Patent Nos. 4.423,143 and 4.560,648 can be

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used, substituting the desired peptide, with or without the use of additional linker groups, for the high molecular weight proteins discussed in the examples provided in these patents, whose complete disclosure is incorporated herein by reference. In the alternative, the specific binding peptide can be produced by recombinant techniques as a fusion protein comprising the specific binding peptide and a second polypeptide, preferably an enzyme label. Most preferably, a functional surrogate of the present invention is incorporated into the primary structure of an enzyme, such as G6PDH or B-galactosidase, in the proximity of its active site. Interaction of the "fusion" enzyme with affinity receptor for the functional surrogate would thus lead to inhibition of the enzyme activity. Hence, DNA constructs comprising DNA sequences encoding an enzyme of choice can be modified by conventional methods to include a DNA insert encoding a functional surrogate. Expressed fusion enzyme can be selected for the desired activity. The inhibition of this activity on exposure of the fusion enzyme to the appropriate affinity receptor is then observed as in the chemically linked combinations of enzyme and functional surrogate. See, e.g., U.S. Patent No. 5,362,625 for representative preparative recombinant techniques involving modified enzymes.

And while the functional surrogates can be used in a wide variety of affinity assays, homogeneous immunoassays would particularly benefit because of the existing inability of such assays to provide accurate, sensitive information regarding macromolecular analytes.

In a preferred embodiment of the invention, a homogeneous fluorescence polarization immunoassay method is provided.

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Fluorescence polarization immunoassay is made possible by the property of fluorescence polarization being affected by the molecular environment. The analyte to be measured (usually a drug, hapten or large molecule in the form of a surrogate) is chemically conjugated to a fluorescent label. A competition is set up between unlabeled analyte in the sample (drug, hapten or large molecule) and the labeled conjugate for limited antibody. The reaction is followed in a Polarization Fluorometer. Using a functional surrogate of the large molecule (or of the drug or hapten for that matter), the fluorescence polarization technique can now be applied as easily and conveniently to the detection of macromolecular analytes. Subsequent combination with an affinity receptor (antibody) alters the molecular environment of the fluorophore due to the presence of a large antibody molecule. Thus, the polarizing property of the fluorophore is altered and monitoring of the reaction and quantitation of analyte can be achieved by following the fluorescence polarization.

More preferably a fluorescent labeled peptide serving as a functional surrogate of a naturally occurring analyte is used and which would compete for limited antibody with the epitope on a macromolecular analyte (supplied as sample).

In a CEDIA format, two fragments (ED and EA) of the enzyme beta-galactosidase are produced by recombinant techniques. Neither of the fragments alone has enzyme activity. When mixed together the 2 fragments combine to form active enzyme. Analyte to be measured is chemically conjugated to the ED fragment, and this conjugated ED fragment can still combine with EA fragment to form active enzyme. However, if ED conjugate is bound by

antibody to analyte then it is unable to combine with EA to form active beta-galactosidase and produce a signal. In this context, "fusion" enzyme fragments comprising ED fragment fused to functional surrogate can also be contemplated.

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Therefore, if a competition is set up between analyte (provided in a sample) and ED conjugate for limited antibody, the presence of analyte in the sample would leave ED available to combine with EA and give active enzyme. Hence, monitoring for beta-galactosidase activity gives a measure of analyte in the sample.

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As already illustrated in the case of EMIT and Fluorescence Polarization, use of small functional surrogate (e.g. a peptide) to conjugate to the ED fragment that would compete for antibody with the epitope on a larger molecule (supplied as sample), would allow for the measurement of larger analytes by CEDIA.

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Accordingly, a method of determining the presence or absence of an analyte of interest in a sample by an affinity assay in accordance with the present invention includes the steps of: (a) providing (i) a labeled conjugate comprising at least one label attached to a functional surrogate capable of competing effectively with an analyte of interest for a limiting amount of an affinity receptor for the analyte, the labeled conjugate exhibiting an activity that is altered on interaction of the labeled conjugate with the affinity receptor and which activity can be measured and related to the presence or absence of the analyte in a given sample, and (ii) the affinity receptor: (b) combining the labeled conjugate and affinity receptor with a sample suspected of containing the analyte to provide a measure of the activity: (c) measuring the activity: and (d) relating the activity to the presence or absence of the analyte in

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the sample. In specific embodiments of the invention this interaction is a binding interaction. Moreover, the functional surrogate may be further characterized as exhibiting a competitive binding profile that is substantially similar to that exhibited by the analyte for the affinity receptor. (See, for example, FIGS. 2, 4, and 6.) In addition, the functional surrogate is further characterized as exhibiting a selective binding affinity (K_a) for the affinity receptor which is substantially similar to that exhibited by the analyte. Hence, while some past work may have used certain octapeptides as a substitute for human chorionic gonadotropin, the binding affinity of the labeled peptide was some two orders of magnitude less than the binding affinity of the natural analyte for anti-hCG.

In a particular method of the invention, step (d), the relating step, comprises comparing the activity with that obtained from at least one control to determine the presence or absence of the analyte in the sample. Preferably, two controls are used, one for a negative result and the second for a positive reading.

As mentioned above, it is preferred that the functional surrogate is obtained by screening a random peptide library with one or more affinity receptors of the analyte.

Most preferably, the random peptide library comprises a plurality of peptides whose structures are not dictated by the primary sequence of the analyte. In specific embodiments, the molecular structure of the functional surrogate may nonetheless correspond to an epitope of the analyte. However, the structure of the epitope may have been previously unknown and would not have been discovered but for the present methods.

And in other cases, the molecular structure of the functional

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surrogate differs from that of a known epitope of the analyte. In this case, the structure of the surrogate may be a composite of a discontinuous epitope or may simply have little or no correlation with the naturally occurring sequence structure. Hence, in certain cases, the molecular structure of the functional surrogate does not include a primary sequence of eight or more continuous amino acid residues which can be found along the naturally occurring sequence of the analyte.

While the functional surrogate of the invention may be of any size suitable for the affinity assay of choice, it preferably has a molecular weight of about 2000 daltons or less. Most preferably, the functional surrogate comprises a peptide of about 1500 daltons or less.

In accordance with the invention, the prescribed combining step may be carried out such that it includes the formation of an affinity receptor-labeled conjugate complex. The combining step may further comprise displacing the labeled conjugate from the complex with the analyte (i.e., a sequential displacement step). Still in other embodiments, the combining step comprises providing competition among the analyte and the labeled conjugate for the affinity receptor. Moreover, the combining step may comprise forming an affinity receptor-analyte complex. Subsequently, the combining step further comprises forming an affinity receptor-labeled conjugate complex. In yet a more specific embodiment of the invention, the combining step comprises (i) mixing the affinity receptor and sample, and (ii) adding the labeled conjugate to the resulting mixture.

Samples suspected of containing an analyte of interest may

be from a wide variety of sources. For example, the sample may be a biological fluid, including but not limited to urine, semen, saliva, sweat, blood, serum, plasma, cerebrospinal fluid, tears, vaginal or nasal fluids. In addition, the sample may be obtained from a cell-free extract.

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In selected embodiments of the invention, the label is selected from the group consisting of a chromogenic gent, a UV absorber, a fluorescent molecule, a chemiluminescent compound, an enzyme, an enzyme fragment, an enzyme substrate, or a group having the potential for exhibiting at least one of the above-recited activities (e.g., after cleavage of a bond). Preferably, the label comprises an enzyme, most preferably one that exhibits glucose-6-phosphate dehydrogenase (G6PDH) activity so that the assay can be performed on a standard clinical chemistry analyzer. If the enzyme has G6PDH activity, then a suitable substrate for the enzyme would include glucose-6-phosphate (G6P) and nicotinamide adenine dinucleotide (NAD). Other suitable enzymes may be those that exhibit lysozyme activity or beta-galactosidase activity.

Various methods of measuring the activity of the labeled conjugate are available depending on the nature of the label. For example, the activity can be measured as a function of the change in the intensity of an absorbance or an emission spectrum, as a function of the change in the polarization or anisotropy of a fluorescence spectrum, as a function of the change in the number of particles observed in a sample mixture, as a function of the change in the amount of a product that is produced by a transformation mediated by the label, as a function of time or a rate of change, to name a few.

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As mentioned above, the analyte can be any molecule of interest, such as a polysaccharide, a polynucleotide, a glycoprotein or a lipid-containing macromolecule. In preferred embodiments, the analyte is a fertility/pregnancy-related hormone, is related to an infectious disease (e.g., a bacterium or a virus), is a cardiac marker or a tumor marker. In still other embodiments, the preferred analytes have already available affinity receptors having a selective, most preferably specific, binding affinity for the analyte, including certain allergens. The molecular weight of the analyte may vary along a wide range, e.g., 200 to 500,000 daltons. Preferably, the analyte has a molecular weight in the range of about 1,000 to about 500,000 daltons, more preferably in the range of about 10,000 to about 200,000 daltons. However, all analytes having a molecular weight in excess of about 100,000 daltons can be detected with the present method.

Thus, the present invention provides, if so desired, a method of determining the presence or absence in a sample of an antibody against an analyte of interest by an affinity assay comprising: (a) providing (i) a labeled conjugate comprising at least one label attached to a functional surrogate capable of competing effectively with an analyte of interest for a limiting amount of an antibody against the analyte, the labeled conjugate exhibiting an activity that is altered on interaction of the labeled conjugate with the antibody and which activity can be measured and related to the presence or absence of the analyte in a given sample; (b) combining the labeled conjugate with a sample suspected of containing the antibody to provide a measure of the activity; (c) measuring the activity; and (d) relating the activity to the presence or absence of the analyte in

the sample.

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In a more specific embodiment, the invention allows a method to be practiced for determining the presence or absence of an analyte of interest in a sample by an affinity assay comprising: (a) providing a labeled conjugate comprising at least one label attached to a functional surrogate capable of competing effectively with an antibody against an analyte of interest for a limiting amount of the analyte, the labeled conjugate exhibiting an activity that is altered on interaction of the labeled conjugate with the analyte or with an affinity receptor for the functional surrogate and which activity can be measured and related to the presence or absence of the analyte in a given sample; (b) combining the labeled conjugate with a sample suspected of containing the analyte to provide a measure of the activity; (c) measuring the activity; and (d) relating the activity to the presence or absence of the analyte in the sample. Preferably, the interaction is at least 50% complete within about 5 minutes of the initiation of the combining step to cut down on incubation times.

In still another embodiment of the invention, a method is enabled for determining the presence or absence of an analyte of interest in a sample by an affinity assay comprising: (a) providing a labeled conjugate comprising at least one label attached to a functional surrogate of an affinity receptor for an analyte of interest, the functional surrogate capable of competing effectively with the affinity receptor for a limiting amount of the analyte, the labeled conjugate exhibiting an activity that is altered on interaction of the labeled conjugate to the analyte and which activity can be measured and related to the amount of the analyte present in a

given sample: (b) combining the labeled conjugate with a sample suspected of containing the analyte to provide a measure of the activity; (c) measuring the activity; and (d) relating the activity to the presence or absence of the analyte in the sample.

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In an enzyme multiplied immunoassay format, the invention provides a method of determining the presence or absence of an analyte of interest in a sample by an homogeneous enzyme affinity assay comprising: (a) providing (i) an enzyme conjugate comprising an enzyme attached to at least one functional surrogate of an analyte of interest, the functional surrogate capable of competing effectively with the analyte for a limiting amount of an affinity receptor for the analyte, the enzyme conjugate exhibiting an activity that is altered on interaction of the enzyme conjugate to the affinity receptor and which activity can be measured and related to the amount of the analyte present in a given sample; (ii) the affinity receptor, and (iii) a substrate for the enzyme: (b) combining the enzyme conjugate, affinity receptor, and enzyme substrate with a sample suspected of containing the analyte to provide a measure of the enzyme activity; (c) measuring the enzyme activity; and (d) relating the enzyme activity to the presence or absence of the analyte in the sample.

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The invention is also applicable in a fluorescence polarization assay in which the presence or absence of an analyte of interest in a sample may be determined by the steps that include:

(a) providing (i) a labeled conjugate comprising a fluorescent material attached to a functional surrogate of an analyte of interest, the functional surrogate capable of competing effectively with an analyte of interest for a limiting amount of an affinity receptor for

the analyte, the labeled conjugate exhibiting an activity that is altered on interaction of the labeled conjugate with the affinity receptor and which activity can be measured and related to the presence or absence of the analyte in a given sample, and (ii) the affinity receptor; (b) combining the labeled conjugate and affinity receptor with a sample suspected of containing the analyte to provide a measure of the activity; (c) measuring the activity; and (d) relating the activity to the presence or absence of the analyte in the sample.

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In a cloned enzyme donor immunoassay format, a method is provided for determining the presence or absence of an analyte of interest in a sample by an homogeneous cloned enzyme donor affinity assay comprising: (a) providing (i) a labeled conjugate comprising an enzyme donor fragment attached to a functional surrogate of an analyte of interest, the functional surrogate capable of competing effectively with the analyte for a limiting amount of an affinity receptor for the analyte, the labeled conjugate, on interaction with an enzyme acceptor fragment, exhibiting an activity that is altered in the presence of the affinity receptor and which activity can be measured and related to the amount of the analyte present in a given sample. (ii) the enzyme acceptor fragment, and (iii) the affinity receptor; (b) combining the labeled conjugate, enzyme acceptor fragment, and affinity receptor with a sample suspected of containing the analyte to provide a measure of the activity; (c) measuring the activity; and (d) relating the activity to the presence or absence of the analyte in the sample.

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An affinity assay kit is also contemplated by the present invention, which comprises: (a) a labeled conjugate diposed in a

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first container means, the labeled conjugate comprising at least one label attached to a functional surrogate of an analyte of interest, the functional surrogate capable of competing effectively with the analyte for a limiting amount of an affinity receptor for the analyte, the labeled conjugate capable of exhibiting an activity that is altered on interaction of the labeled conjugate with the affinity receptor and which activity can be measured and related to the amount of the analyte present in a given sample; and (b) disposed in a second container means the affinity receptor and, optionally, any substance required for the labeled conjugate to exhibit the activity, such as an enzyme substrate or an enzyme acceptor fragment.

Most importantly, in a particular embodiment of the present invention, a functional surrogate of an analyte of interest is contemplated which comprises a peptide having an immunoreactive group that allows the surrogate to compete effectively with the analyte for a limiting amount of an affinity receptor for the analyte.

It has been discovered, for instance, that certain binding peptides listed in Tables 6-16, include motifs that appear to be important to selective binding affinity. Certain of the sequences flanking the motifs may also be necessary in some cases.

Specifically, for ferritin, such motifs may include: AGRALFH (SEQ ID NO:80), HGRAMFQ (SEQ ID NO:81), GGQAMFN (SEQ ID NO:82), GGSAMFS (SEQ ID NO:83), GGEALFK (SEQ ID NO:84), GGRSLFQ (SEQ ID NO:85), GGMSLFQ (SEQ ID NO:86), GGASLFQ (SEQ ID NO:87), IGASLFQ (SEQ ID NO:88), SSSALFQ (SEQ ID NO:89), SNSALFQ (SEQ ID NO:90), PQRAFFQ (SEQ ID NO:91), SINPT (SEQ ID NO:92), SINGTP (SEQ ID NO:93), GGDALFT (SEQ ID NO:94), SGGSSFW (SEQ

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ID NO:95), GNTVMFQ (SEQ ID NO:96), FCGAMFC (SEQ ID NO:97), SKDSFFQ (SEQ ID NO:98), PASAFFQ (SEQ ID NO:99), HSSSLFQ (SEQ ID NO:100), NGSSLFN (SEQ ID NO:101), GGRAFFL (SEQ ID NO:102), AGRAFFR (SEQ ID NO:103), SQSS FQ (SEQ ID NO:104), HSSSLF (SEQ ID NO:105), HSSSLFQ (SEQ ID NO:106), AGAPLFQ (SEQ ID NO:107), RGNAFFK (SEQ ID NO:108), GGEVLFK (SEQ ID NO:109), GGSAAFQ (SEQ ID NO:110), GGEALFQ (SEQ ID NO:111), GGRALFA (SEQ ID NO:112), RVSTLFQ (SEQ ID NO:113), AGLALFQ (SEQ ID NO:114), HSSSFFQ (SEQ ID NO:115), SSSAFFQ (SEQ ID NO:116), PITNMFQ (SEQ ID NO:117), AGRAFFR (SEQ ID NO:118), GGDALFT (SEQ ID NO:119), GGHSFFK (SEQ ID NO:120), GGMSLFQ (SEQ ID NO:121), SGSSMFQ (SEQ ID NO:122), SSSSLFQ (SEQ ID NO:123). HSSSLFQ (SEQ ID NO:124), CRGSLFC (SEQ ID NO:125). *GGMALFP (SEQ ID NO:126), GGGAMFQ (SEQ ID NO:127), RGRAMFK (SEQ ID NO:128), HSSSMFQ (SEQ ID NO:129), GGRSLFT (SEQ ID NO:130), GGASLFL (SEQ ID NO:131) or GARALFL (SEQ ID NO:132).

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For the hepatitis antigen (e.g., hepatitis A. B or C, but especially B), certain motifs include HPLY (SEQ ID NO:133), HPIY (SEQ ID NO:134), GPPHL (SEQ ID NO:135), GPGPL (SEQ ID NO:136), GPGHL (SEQ ID NO:137), GPRHL (SEQ ID NO:138), VPPHL (SEQ ID NO:139), PPAHL (SEQ ID NO:140), PPPNL (SEQ ID NO:141). ARSDE (SEQ ID NO:142), LRSRE (SEQ ID NO:143), LRSAE (SEQ ID NO:144), KTVLPR (SEQ ID NO:145), GEVLPK (SEQ ID NO:146), GAVLPR (SEQ ID NO:147), GAVLAK (SEQ ID NO:148), GPKHL (SEQ ID NO:147), GAVLAK (SEQ ID NO:148), GPKHL (SEQ ID

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NO:149), GPDHL (SEQ ID NO:150), GPEHL (SEQ ID NO:151), STSSIGPLR (SEQ ID NO:152), SNTPRGPLK (SEQ ID NO:153), STTSAGPRK (SEQ ID NO:154), SGTARGPTK (SEQ ID NO:155), SLTSSGPIK (SEQ ID NO:156), RCPSDGNCY (SEQ ID NO:157) or RCPSDGLCY (SEQ ID NO:158). Accordingly, preferred binding peptides are those that include at least the primary sequence motifs depicted in Tables 1-2 and 6-16.

Furthermore, consensus sequences can be inferred from the amino acid sequences depicted in Tables 6-16. Such consensus sequences may have a particular residue conserved at a particular position. At other positions, the amino acids may vary within a particular type of residue, including but not limited to, hydrophobic amino acids (such as A, V, L, I, P, F and the like -- symbol Φ), hydrophilic residues (e.g., S, T, K, R, H, D, E, C and the like -- symbol Ψ), basic residues (e.g., K, R, H -- symbol θ), acidic residues (e.g., D, E -- symbol σ), aromatic residues (such as F, Y, W, H and the like -- symbol π) or amide containing residues (e.g., N, Q -- symbol Ω). Some residues, such as G, C or M may be considered either hydrophobic or hydrophilic. The symbol X means that a position is not conserved and may include any residue.

Hence, for a surrogate for hepatitis B antigen, certain consensus sequences can be identified, including HP(I/L)Y (SEQ ID NO:159), (SEQ ID NO:160), GPXHL (SEQ ID NO:161), (A/L)RSXE (SEQ ID NO:162), (SEQ ID NO:163), GXVLP0 (SEQ ID NO:164), STTXXGPXK (SEQ ID NO:165) or CPSDGNCY (SEQ ID NO:166). Possible consensus sequences for ferritin antigen surrogates may include GGX(A/S)LFQ (SEQ ID NO:167), (SEQ ID NO:168), SIN(P/G)TP (SEQ ID NO:169), (SEQ ID

NO:170) or GGMALFP (SEQ ID NO:171). It is important to point out, however, that other consensus sequences can be gleaned from the sequences presented in the Tables herein. Such sequences are, of course, considered part of this invention.

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In addition to the labeled conjugates, the present invention also contemplates recombinant DNA constructs which comprise DNA sequences encoding a functional surrogate of an analyte of interest, the functional surrogate capable of competing effectively with the analyte for a limiting amount of an affinity receptor for the analyte. In particular, constructs comprising DNA sequences, as depicted in Tables 6-16, are particularly desired. Preferred sequences will be at least those that encode a primary sequence motif of the present invention.

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Transforming vectors or expression vehicles including these constructs are also contemplated, as well as bacteriophage and viable eucaryotic and procaryotic cells transformed with such vectors or vehicles. Microorganisms can, of course, be infected with the selected bacteriophage, resulting in expression of the encoded peptides. For the production of large quantities of peptide or fusion proteins including the peptides, yeast vectors can be constructed which direct the secretion of encoded peptides into the culture medium. (See, for example, U. S. Patent No. 4.546,082, the disclosure of which is incorporated by reference herein.)

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It has thus been discovered that functional surrogates of naturally occurring analytes can be obtained by a method that includes the steps of: (a) selecting an affinity receptor exhibiting a selective affinity for an analyte of interest; (b) screening a random peptide library with the affinity receptor for a binding peptide; (c)

isolating the binding peptide and identifying its structure. The peptide isolated and identified can then be synthesized and its capacity to compete with the analyte for a limiting amount of the affinity receptor verified. As discussed above, the use of a phage displayed random peptide library is particularly preferred.

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The preparation and characterization of the preferred phagedisplayed random peptide libraries have been described elsewhere. See, for example, Kay, B.K. et al. in Gene (1993) 128:59-65 and International Application No. PCT/US94/0977, for a description of the preparation of phage-displayed random peptide libraries. For a description of the libraries known as R8C, D38, and DC43, see below. In particular, by cloning degenerate oligonucleotides of fixed length into bacteriophage vectors, recombinant libraries of random peptides can be generated which are expressed at the amino-terminus of the pIII protein on the surface of M13 viral particles. (There are 3-5 copies of the pIII-fusion on the surface of each particle.) Phage display offers several conveniences: first, the expressed peptides are on the surface of the viral particles and accessible for interactions; second, the recombinant viral particles are stable (i.e., can be frozen, exposed to pH extremes); third, the viruses can be amplified; and fourth, each viral particle contains the DNA encoding the recombinant genome. Consequently, these libraries can be screened by isolating viral particles that bind to targets. The isolates can be grown up overnight, and the displayed peptide sequence responsible for binding can be deduced by DNA sequencing.

These libraries have approximately >10⁸ different recombinants, and nucleotide sequencing of the inserts suggests that

the expressed peptides are indeed random in amino acid sequence.

6. <u>Examples</u>

The following Examples are provided to assist the reader further, which Examples describe selected materials, compositions, and methods for use in particular embodiments and which are illustrative of the invention, as a whole.

Examples of materials used in the methods of the invention are set forth below.

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Chemical Reagents

10xPBS; Dulbecco's PBS x 10 (JRH Biosciences, Lenexa, KS, Cat # 59331-78P)

PBS; phosphate buffered saline (1-10 dilution in water of

Dulbecco's 10x PBS)

BSA; bovine serum albumin (Sigma, St Louis, MO; Cat # A7906) Tween 20; polyoxyethylene sorbitan monolaurate (Sigma, St Louis, MO; Cat # P1379)

PBS/BSA; PBS with 1% BSA

20 PBT; PBS with BSA (1%) & Tween 20 (0.05%)

X-Gal; X-Gal (Jersey Lab. Supply; Livingston, NJ; Cat # X266)

DMF; dimethyl formamide (Sigma, St Louis, MO; Cat # D4254)

X-Gal solution; 2% X-Gal in DMF

TMB; tetramethyl benzidine substrate (KPL, Gaithersburg, MD; Cat # 50-76-00)

- - -

2xYT broth;

Streptavidin-coated microtiter plates: Reactibind (Pierce, Rockford, IL, Cat # 15120)

Microtiter plates (Immulon 4, Dynatech, Chantilly, VA; Cat # 011-010-3855)

DH5xF': E. coli cells

IPTG; isopropyl-\u00a3-D-thiogalactopyranosidase (Jersey Lab Supply,

Livingston, NJ;

Cat # 1555)

IPTG solution; 100 mM IPTG in water

SM buffer

G6PDH; glucose 6 phosphate dehydrogenase (Sigma, St. Louis,

10 MO; Cat # G5760)

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NHS; N-hydroxysuccinimide (Sigma, St. Louis, MO; Cat # H7377)

DCC; dicyclohexylcarbodiimide (Sigma, St. Louis, MO: Cat #

D3128)

DMSO; dimethylsulfoxide (Sigma, St. Louis MO; Cat # D5879)

15 Tris; TRIZMA base (Sigma, St Louis, MO; Cat # T8524)

G6P: glucose 6 phosphate, sodium salt (Sigma, St. Louis, MO; Cat

G7879)

Carbitol: diethyleneglycol monoethylether (Sigma, St. Louis, MO.;

Cat # D1265)

20 NADH: nicotinamide adenine dinucleotide, reduced form (Sigma,

St. Louis, MO; Cat # N6005)

PEG; polyethylene glycol ave. MW 8000 (Sigma, St. Louis, MO;

Cat # P2139)

NaCl; sodium chloride (Sigma, St. Louis, Mo.; Cat # S9625)

25 PEG/NaCl; 20% PEG 8000 in 2.5 M NaCl

Immuno Reagents

Affinity purified goat anti-HBsAg (OEM Concepts, Toms River,

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NJ; Cat # G5-V18)

Affinity purified sheep anti-ferritin (The Binding Site, San Diego, CA; Cat # AU055)

Monoclonal mouse anti-HBsAg (OEM Concepts, Toms River, NJ;

5 Cat # M2-V18)

Goat anti-mouse IgG HRP conjugate (OEM Concepts, Toms River, NJ.;

Cat # G5-MG16-2)

Rabbit anti-goat IgG HRP conjugate (OEM Concepts, Toms River, NJ:

Cat # R5-GG10-2)

Rabbit anti-sheep IgG HRP conjugate (OEM Concepts, Toms River, NJ;

Cat #R5-SG10-2)

Normal non-immune mouse IgG (OEM Concepts, Toms River, NJ: Cat # M6-G10)

Normal non-immune goat IgG (OEM Concepts, Toms River, NJ; Cat # G6-G10)

Normal non-immune sheep IgG (OEM Concepts, Toms River, NJ; Cat # S8-G10)

rHBsAg; recombinant HBsAg AY antigen (OEM Concepts, Toms River, NJ;

Cat # H7-V57)

Ferritin antigen (OEM Concepts, Toms River, NJ; Cat # H6-MO5)

Rabbit anti-M13 HRP conjugate (Pharmacia, Piscataway, NJ, Cat # 27-9402A)

Phage Display Libraries

The following Libraries were used for biopanning with mouse anti-HBsAg and goat anti-HBsAg: R8C and R26.

For biopanning with sheep anti-ferritin, the following libraries were used: D38; DC43; and R26.

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6.1. <u>Preparation of a Phage Display Random Peptide</u> <u>Library</u>

6.1.1. <u>General Synthesis and Assembly of Oligonucleotides</u>

Random sequence oligonucleotide inserts flanked by selected cloning sites were synthesized with an applied Biosystems 380a synthesizer (Foster City, CA), and the full-length oligonucleotides were purified by HPLC.

Five micrograms of each of the pair of oligonucleotides were mixed together in buffer (10 mM Tris-HCl, pH 8.3, 15 mM KCl, 0.001% gelatin, 1.5 mM magnesium chloride), with 0.1 % Triton X-100, 2 mM dNTP's, and 20 units of Tag DNA polymerase. The assembly reaction mixtures were incubated at 72 °C for 30 seconds and then 30 °C for 30 seconds; this cycle was repeated 60 times. It should be noted that the assembly reaction is not PCR, since a denaturation step was not used. Fill-in reactions were carried out in a thermal cycling, device (Ericomp, LaJolla, CA) with the following protocol: 30 seconds at 72 °C, 30 seconds at 30 °C, repeated for 60 cycles. The lower temperature allows for annealing of the six base complementary region between the two sets of the oligonucleotide pairs. The reaction products were phenol/chloroform extracted and ethanol precipitated. Greater than 90% of the nucleotides were found to have been converted to double stranded synthetic oligonucleotides.

After resuspension in 300 µL of buffer containing 10 mM Tris-HCI, pH 7.5, 1 mM EDTA (TE buffer), the ends of the oligonucleotide fragments were cleaved with Xba I and Xho I (New England BioLabs, Beverly, MA) according to the supplier's

recommendations. The fragments were purified by 4% agarose gel electrophoresis. The band of correct size was removed and electroeluted, concentrated by ethanol precipitation and resuspended in 100 µL TE buffer. Approximately 5% of the assembled oligonucleotides can be expected to have internal Xho I or Xba I sites; however, only the full-length molecules were used in the ligation step of the assembly scheme. The concentration of the synthetic oligonucleotide fragments was estimated by comparing the intensity on an ethidium bromide stained gel run along with appropriate quantitated markers. All DNA manipulations not described in detail were performed according to Sambrook, et al., infra.

To demonstrate that the assembled enzyme digested oligonucleotides could be ligated, the synthesized DNA fragments were examined for their ability to self-ligate. The digested fragments were incubated overnight at 18 °C in ligation buffer with T4 DNA ligase. When the ligation products were examined by agarose gel electrophoresis, a concatamer of bands was visible upon ethidium bromide staining. As many as five different unit length concatamer bands (i.e., dimer, trimer, tetramer, pentamer, hexamer) were evident, suggesting that the synthesized DNA fragments were efficient substrates for ligation.

6.1.2. Construction of Vectors

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The construction of the M13 derived phage vectors useful for expressing a random peptide library has been recently described (Fowlkes, D. et al. *BioTech.* (1992) 13:422-427). To express the library, an M13 derived vector, m663, was constructed

as described in Fowlkes. The m663 vector contains the pIII gene having a c-myc-epitope, i.e., as a stuffer fragment, introduced at the mature N-terminal end, flanked by Xho I and Xba I restriction sites (See also, Figure I of Fowlkes).

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6.1.3. Expression of the Random Pentide Library

The synthesized oligonucleotides were then ligated to Xho I and Xba I double-digested m663 RF DNA containing the pIII gene (Fowlkes) by incubation with ligase overnight at 12 °C. More particularly, 50 ng of vector DNA and 5 ng of the digested synthesized DNA and were mixed together in 50 µL ligation buffer (50 mM Tris, pH 8.0, 10 mM MgCl₂, 20 mM DTT, 0.1 mM ATP) with T4 DNA ligase. After overnight ligation at 12 °C, the DNA was concentrated by ethanol precipitation and washed with 70% ethanol. The ligated DNA was then introduced into E. coli (DH5 α F'; GIBCO BRL, Gaithersburg, MD) by electroporation.

A small aliquot of the electroporated cells was plated and the number of plaques counted to determine that 10⁸ recombinants were generated. The library of *E. coli* cells containing recombinant vectors was plated at a high density (~400,000 per 150 mM petri plate) for a single amplification of the recombinant phage. After 8 hr, the recombinant bacteriophage were recovered by washing each plate for 18 hr with SMG buffer (100 mM NaCl. 10 mM Tris-HCl, pH 7.5, 10 mM MgCl₂, 0.05% gelatin) and after the addition of glycerol to 50% were frozen at -80 °C. The library thus formed had a working titer of ~2 x 10¹¹ pfu/ml.

6.2. Preparation of R8C Library

Referring now to Figure 7, two oligonucleotides were synthesized on an Applied Biosystems Model 380a machine with the sequence 5'-

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TGACGTCTCGAGTTGTNNKNNKNNKNNKNNKNNKNNKNN KTGTGGATCTAGAAGGATC-3' (SEQ ID NO:172) and 5'-GATCCTTCTAGATCC-3" (SEQ ID NO:173), where N is an equimolar ratio of deoxynucleotides A. C, G, and T, and K is an

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equimolar ratio of G and T. Fifty pmol of each oligonucleotide was incubated at 42 °C for 5 min, then 37 °C for 15 minutes in 50 μL of Sequenase TM buffer (U.S. Biochemicals, Cleveland, OH) with

0.1 μg/μL acetylated BSA, and 10 mM DTT. After annealing, 10 units of Sequenase TM (U.S. Biochemicals) and 0.2 mM of each

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dNTP were added and incubated at 37 °C for 15 min. The sample was then heated at 65 °C for 2 hr, digested with 100 units of both

Xho I and Xba I (New England BioLabs, Beverly, MA), phenol extracted, ethanol precipitated, and resolved on a 15% non-

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denaturing polyacrylamide gel. The assembled, digested fragment was gel purified prior to ligation. The vector, m663 (Fowlkes, D.

et al. Biotech. (1992) 13:422-427), was prepared by digestion with Xho I and Xba I, calf alkaline phosphatase (Boehringer Mannheim,

Indianapolis, IN) treatment, phenol extracted, and purified by agarose gel electrophoresis. To ligate, 20 μg vector was combined

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with 0.2 µg insert in 3 mL with T4 DNA ligase (Boehringer Mannheim), according to the manufacturer. After removal of the protein and buffer by phenol extraction and ethanol precipitation,

the ligated DNA was electroporated into XL1-Blue E. coli (Stratagene, San Diego, CA) and plated for eight hours at 37 °C.

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To recover the recombinant phage, the top agar was collected with a spatula, mixed with an equal volume of 100 mM NaCl, 10 mM MgCl₂, and 50 mM Tris-HCl (pH 7.5), and disrupted by two passes through an 18-gauge syringe needle. The bacterial cells were removed by centrifugation, and phage particles were collected by polyethylene glycol precipitation and stored at -70 °C in 25% glycerol. The library had 10⁸ total recombinants and a working titer of 6 x 10¹³ pfu/mL.

Members of the library were checked for inserts by the polymerase chain reaction (Saiki, et al. Science (1988) 239:487-491). Individual plaques on a petri plate were touched with a sterile toothpick and the tip was stirred into 2xYT with F* E. coli bacteria and incubated overnight at 37 °C with aeration. Five microliters of the phage supernatant were then transferred to new tubes containing buffer (67 mM Tris-HCl, pH 8.8/10 mM β -mercaptoethanol/16.6 mM ammonium sulfate/6.7 mM EDTA/50 μg bovine serum albumin per mL), 0.1 mM deoxynucleotide triphosphates, and 1.25 units of Taq DNA polymerase (Boehringer Mannheim, Indianapolis, IN) with 100 pmoles of oligonucleotide primers. The primers flanked the cloning site in the pIII gene of m663 (5'-TTCACCTCGAAAGCAAGCTG-3' (SEQ ID NO:174) and 5'-CCTCATAGTTAGCGTAACG-3' (SEQ ID NO:175)). The assembly reactions were incubated at 94 °C for 1 min. 56 °C for 2 min, and 72 °C for 3 min; this cycle was repeated 24 times. The reaction products were then resolved by electrophoresis on a NuSieve 2.0% agarose gel (FMC, Rockland, ME). Gels revealed that for 20 plaques tested, all were recombinant and had single inserts of the expected size.

The R26, D38, and DC43 libraries were prepared similarly based on the schematic provided in FIGS, 8, 9, and 10.

6.3. Biopanning

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Biopanning was carried out on microtiter wells that had been coated with 100μL of immune antiserum (e.g., mouse monoclonal anti-HBsAg) diluted in PBS to 5 ug/mL. The wells were blocked with 100 μL PBS/BSA overnight at 4 °C, washed 3x with 200 μL PBS/BSA, then banged dry on paper towels.

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Phage libraries to be panned were diluted in PBS to approximately 10^{10} to 10^{11} pfu per 40 μ L. Diluted phage library (40 μ L) was added to each well, the plate covered, and allowed to incubate on a rocker for 1 hr at room temperature.

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The contents of the wells were dumped, and the wells washed 10x with $200~\mu L$ PBS/BSA, then banged dry on paper towels. Bound phage was then eluted by adding $50~\mu L$ of 0.05~M glycine pH 2.0 to each well and incubating for 5~min at room temperature on the rocker.

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 $50~\mu L$ of 0.2 M Phosphate pH 7.6 was added to a separate tube and, after the 5 minute incubation, the $50~\mu L$ of eluted phage (in the glycine pH 2.0) was added to the 0.2 M phosphate pH 7.6.

6.4. <u>Isolation and Characterization of Antibody Binding</u> Phage from Libraries

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To illustrate the invention, antibody (monoclonal mouse anti-HBsAg or affinity purified polyclonal goat anti-HBsAg or affinity purified sheep anti-human ferritin) was used to biopan the libraries. Three (3) rounds of biopanning were performed, and

binders were amplified once between the 1st and 2nd rounds.

Binders from the 3rd round of panning were grown on agar plates. Routinely 30 individual plaques were picked manually, grown in liquid culture overnight at 37 °C and assayed in the M13 phage ELISA to determine specific binding to antibody.

DNA from positive clones was isolated and sequenced. The sequences were then examined manually and any consensus motifs determined.

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6.5. ELISA for Phage M13

Specific phage binders were confirmed with an ELISA for M13. The assay was performed on phage cultures that had been grown overnight from the 30 plaques that had been picked individually after the 3rd round of biopanning.

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Alternate rows of a microtiter plate were coated with 50 μ L of immune antibody at 5 μ g/mL in PBS (e.g., mouse anti HBsAg), appropriate non-immune IgG at 5 μ g/mL in PBS, and PBS/BSA for one hour at room temperature. The solutions were dumped and the wells blocked for a minimum of one hour at room temperature with 200 μ L PBS/BSA. The solutions were dumped, the wells washed 3x with 200 μ L PBS/BSA, and then banged dry on paper towels.

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Phage to be assayed (phage growth medium supernatant from a 6h liquid growth) was diluted 1-100 in PBS/BSA, and 100 μL of this dilution was added to each of 6 wells - 2 coated with immune IgG, 2 coated with non-immune IgG, and 2 coated with BSA; i.e., each phage sample was assayed in duplicate, with non-specific binding (NSB) controls of non-immune serum and BSA.

William Balletin

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The plate was incubated at room temperature for 1 hour,

then washed 3x with $200~\mu L$ PBS/BSA and banged dry on paper towels. $100~\mu L$ anti-M13 HRP (diluted 1:8000 in PBS/BSA) was added to each well, the plate incubated for 1 hour at room temperature, then washed 3x with $200~\mu L$ PBS/BSA and banged dry on paper towels.

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 $100~\mu L$ TMB substrate was added to each well and the blue color allowed to develop for about 5 minutes. The plate was then read at 620 nm on the SLT 340 ATTC or Magnetic Devices V max Microplate Reader.

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Positive phage were identified as binding to the antibody coated wells, but not the wells coated with non-immune IgG or BSA.

6.6. Phage Amplification

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Phage amplification on plates and in liquid was carried out as described, McConnell, SJ., Uveges, AJ., & Spinella, DG. *Biotechniques* (1995) 18:803-806.

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When amplification was used in between rounds 1 and 2 of biopanning, debris was removed from solution by centrifugation at 1000 rpm for 10 minutes at 4 °C, then phage was precipitated with 0.2 vol PEG/NaCl for 2h on ice. Supernatant was removed after centrifugation at 10,000 rpm for 15 minutes at 4 °C and phage dissolved in 100 μ L of PBS/BSA and transferred to a microtube. The solution was clarified by centrifugation (10,000 rpm, 10 minutes) and used immediately.

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6.7. DNA Sequencing

DNA sequencing of phage ciones confirmed positive from

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the M13 ELISA was used to determine the amino acid sequence of the random peptide insert.

Single stranded DNA template was prepared using the Dynabeads R lacZ ssM13 Purification Kit (Applied Biosystems/Perkin Elmer, Foster City, CA; Cat # 401436) according to the protocol provided by the manufacturer.

DNA sequencing was performed on a model 373A DNA Sequencer (Applied Biosystems, Inc., Foster City, CA) using the Prism ™ Ready Reaction DyeDeoxy Terminator Cycle Sequencing Kit (Applied Biosystems/Perkin Elmer, Foster City, CA; Cat # 401384) according to the protocol provided by the manufacturer.

PC/GENE Release 6.7 (Intelligenetics, Mountain View, CA) was used to analyze the DNA sequence, and determine the deduced amino acid sequence of the random peptide insert.

6.7.1. Anti-HBsAg Binders Mouse Monoclonal Anti-HBsAg

Libraries giving positive phage binders when panned with monoclonal anti-HBsAg were R26 and R8C. The sequences and consensus motifs are shown in Table 4. From a total of 26 unique (non-sibling) sequences examined, four consensus motifs could be recognized, with only three sequences not containing any of the four motifs. None of the motifs appeared to match any portion of the HBsAg primary sequence, hence these motifs were regarded as mimetopes.

6.7.2. <u>Affinity Purified Goat Polyclonal Anti-HBsAg</u> Libraries giving positive phage binders when panned

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with goat anti-HBsAg were R26a, b, c & d, and R8C. The sequences and consensus motifs are shown in the Table 4. From a total of 12 unique (non-sibling) sequences examined, two consensus motifs could be recognized. The four M13 ELISA positive phage from the R8C Library failed to give any consensus motif that could be recognized.

Both of the motifs appeared to match a portion of the HBsAg primary sequence, hence these motifs were regarded as epitopes. Only the R8C Library failed to produce any consensus motif from the M13 ELISA confirmed positive binders.

6.8. Solid Phase Peptide Synthesis

Consensus peptide motifs deduced from DNA sequencing were assembled using a modification of Merrifield's solid phase method (Merrifield, 1963) using either standard HBTU chemistry on a Model 430A Peptide Synthesizer (Perkin Elmer/Applied Biosystems, Foster City, CA) or using the same chemistry on a Symphony Multiple Peptide Synthesizer (Rainin/PTI, Woburn, MA) Resin used for assembly was Tentagel S RAM (Tubingen, Germany).

9-Fluorenylmethoxycarbonyl (Fmoc) derivatives of amino acids were used throughout, with side chains blocked by t-butyl type moieties. The same HBTU chemistry was used to add biotin at the amino terminus of some peptides. Stepwise reaction efficiencies were monitored by ninhydrin (Kaiser, 1970), and were typically >95%.

Resin cleavage and side chain deblocking were performed simultaneously for 120 minutes at room temperature using reagent

K (King, 1990). Multiple washes with t-butyl methyl ether were performed after cleavage to remove scavengers.

Peptides were desalted and purified via preparative reverse phase HPLC before lyophilization and storage at room temperature. Amino acid analyses (AAA) were carried out on a Beckman System Gold (Beckman, Fullerton, CA) after vapor phase HCl hydrolysis (Meltzer, 1987). Mass Spectroscopy (FAB-MS) was performed by M Scan (West Chester, PA.).

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6.9. Oxidation of Pentides

Peptides designed to contain a disulfide bond were oxidized immediately. Crude peptide was dissolved at 0.5 mg/ml in 50 mM sodium phosphate, pH 8.0, with 1% acetonitrile added as antimicrobial. The mixture was stoppered loosely with glass wool and allowed to stir gently on a magnetic stirrer in contact with air. Oxidation was allowed to proceed for 12-36 hours, and the reaction was monitored with 5,5 dithio-bis(nitrobenzoic acid) (Deakin, 1963) to determine the end-point.

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6.10. Titration of Peptide and Primary Ab by ELISA

Reaction of synthesized peptides with antibody was detected using a microtiter plate based ELISA. Varying amounts of synthetic peptide were immobilized either via attached biotin (to streptavidin coated plates) or chemically. Binding of varying amounts of added antibody to the immobilized synthetic peptide was detected with HRP conjugated 2nd antibody. As shown in FIG. 1, the more synthetic peptide is present in the microtiter plate, the greater the response from added antibody reagents. Hence, the

synthetic peptides of the present invention exhibit the selective binding characteristics expected of a functional surrogate for naturally occurring hepatitis B surface antigen. Similar results are shown in FIGS. 3 and 5.

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6.10.1 <u>Immobilization of Biotinylated Peptides</u>

Biotinylated peptide was diluted in PBT and coated onto streptavidin plates at 4°C overnight. 100µL of diluted solution was used per well. The contents were dumped, the plate washed 3x with 200 µL PBS/BSA/T, and banged dry on paper towels.

6.10.2. <u>Immobilization of Non-Biotinylated</u> Peptides With Carbodiimide

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Non-biotinylated peptides were immobilized onto microtiter plates using CDI by the method of Dagensis, P., in *Anal. Biochm.* (1994) 222:149-156. Irrespective of the immobilization procedure, the subsequent assay was carried out as follows:

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6.10.3. Assay Procedure

Primary antibody (e.g., mouse anti-HBsAg) was diluted in PBT. 100 μ L of each dilution was added to appropriate wells and incubated at room temperature for one hour on a rocker. The contents were dumped, the plate washed 3x with 200 μ L PBS/BSA/T, and banged dry on paper towels.

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Appropriate 2nd Ab HRP conjugate (e.g., goat anti-Mouse HRP) was diluted 1:15,000 in PBT, 100 μ L added to each well, and incubated for one hour at room temperature on a rocker. The contents were dumped, the plate washed 3x with 200 μ L

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PBS/BSA/T, and banged dry on paper towels.

 $100~\mu L$ of TMB substrate was added to each well and allow to develop for 15-30 minutes at room temperature. The plate was then read at 650nm on the SLT 340 ATTC or Magnetic Devices V max Microplate Reader.

6.11. ELISAs for Antigen Using Immobilized Antigen or Binding Peptide

The ability of a synthetic binding peptide to act as a functional surrogate of the natural antigen was demonstrated by constructing competitive and sequential ELISAs for antigen using either immobilized antigen or synthetic peptide bound to a solid phase. The resulting competitive binding profiles for three selected synthetic peptides are presented in FIGS. 2, 4 and 6. As illustrated in these figures, the functional surrogates of the present invention can compete effectively for limited antibody in the presence of natural whole antigen.

6.11.1. Immobilization of Antigen, Biotinylated and Non-Biotinylated Peptides

Whole antigen was diluted in PBS and immobilized onto microtiter plates using passive adsorption at 4 °C overnight. Biotinylated peptides were immobilized on streptavidin coated plates, and non-biotinylate peptides were covalently bound using CDI. as previously described.

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6.11.2. Sequential ELISA

Antigen and antibody were diluted in PBT. Equal volumes of each antigen concentration and antibody solution were mixed in separate glass 12 x 75 mm tubes and incubated at room temperature for 90 minutes. Triplicate 100 µL portions of each mixture were added to appropriate wells of a microtiter plate and incubated at room temperature for 2.5h on a rocker.

The subsequent procedure was used irrespective of immobilization method or whether the assay was competitive or sequential.

The contents were dumped, the plate washed 3x with 200 μ L PBS/BSA/T, and banged dry on paper towels.

The appropriate second Ab HRP conjugate was diluted 1:15000 in PBT, and 100 μ L of the diluted solution was added to each well. The resulting mixture was incubated for one hour at room temperature on a rocker. The contents were the dumped, the plate washed 3x with 200 μ L PBS/BSA/T, and banged dry on paper towels.

 $100~\mu L$ of TMB substrate was added to each well and the reaction was allowed to develop for 30 minutes at room temperature. The plate was then read at 650 nm on the SLT 340 ATTC or Magnetic Devices V max Microplate Reader.

6.12. Conjugation of Peptide to G6PDH

The peptide N-hydroxysucccinimide ester (0.1 M) was prepared in dimethyl formamide with equimolar concentrations of NHS (0.1 M) and DCC (0.1 M). The active ester was then coupled to G6PDH as described by Oellerich (1986).

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The functional surrogates are normally bonded to the enzyme directly through a single or double bond, or indirectly through a linking group. The functional groups of the enzyme which are available for linking are amino (including guanidino), hydroxy, carboxyi, mercapto, and activated aromatic groups of imidazole.

The binding peptides have a great diversity of functional groups available for coupling; additionally, routine modifications of these functionalities may be made to facilitate the conjugation, e.g., conversion of keto to hydroxy, or olefin to aldehyde or carboxylic acid.

Where a linking group is employed for bonding the functional surrogate to the enzyme label, normally the linker will be attached to the peptide to provide a means for coupling of the peptide to the enzyme. This conjugation may be achieved in a single step or may require multiple steps, including blocking and unblocking of active sites of the peptide other than those involved in providing the linking group.

Where the enzyme is to be linked through a carboxyl group of the functional surrogate or a linker bonded to the functional surrogate, either esters or amides will be prepared. The functional surrogate may be bonded to any of the linking groups which are appropriate to provide a link between the functional surrogate and the alcohol or amine group of the enzyme to form the ester or amide group, respectively. When the enzyme has an activated aromatic ring, the functional surrogate may be bonded to an aromatic diazonium salt to provide the desired bridge.

When bonding a functional surrogate through a linking

group to an enzyme, the bonds formed must be stable under assay conditions and the conditions used for carrying out the coupling reactions must not result in an inactive enzyme conjugate.

Additionally, the enzyme must not prevent binding of receptor to functional surrogate.

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For instance, if the functional surrogate has an amino group the amino may be derivatized to alpha-bromoacetamide. This group can then form a C-N bond to an amino acid of an enzyme that has a free amino group (such as lysine). If the functional surrogate has a keto group, the carbonyl may be condensed directly with an amine group of the enzyme, or the O-carboxy methyloxime may be prepared with O-carboxymethyl hydroxylamine. A mixed anhydride, with isobutyl chloroformate is formed, which can then form the carboxamide with the amino group of a lysine. Where a carboxyl group is present in the functional surrogate, this group may be derivatized to the monoamide of phenylenediamine. The intermediate can then be diazotized to form a diazo salt suitable for coupling to a tyrosine in the enzyme.

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Also a hydroxyl group is present in the functional surrogate it can be reacted with succinic anhydride to form a monoester. The free carboxy group can then be used to prepare the mixed anhydride, which in turn can be reacted with an amino group in the enzyme.

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Where an amino group is present in the functional surrogate this may be reacted with maleic anhydride to give the maleimide. The maleimide may then be reacted with cysteine in the enzyme to give a 3-thiosuccinimide.

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6.13. Procedure for EMIT Assay

Using a set amount of conjugate, a solution of antibody is titrated to give approximately 40-60% inhibition of activity. In the presence of antigen, the inhibitory effect of added conjugate on antibody activity is proportional to the amount of antigen present. Thus, a dose response curve of G6PDH activity vs antigen concentration can be obtained.

6.14 Procedure for Testing Peptide-G6PDH Conjugate

G6PDH activity is measured by the rate of conversion of NAD to NADH as measured by increase in absorbance at 340 nm at 37 °C. The dilution of conjugate for use in the EMIT assay is determined from the results.

Peptide-G6PDH conjugate (conjugate) is diluted 1:10, 20, 40, 80, 160, 320 and 640 in a buffer of 0.218 M Tris pH 8.0 containing 1 g/L BSA and 7g/L Glucose 6 phosphate.

25 μL portions of diluted conjugate are added to appropriate wells of a microtiter plate. 100 μL of buffer (0.013 M Tris pH 6.0 containing l g/L BSA, 20 g/L NaCl, l mL/L Tween 20) is added to each well and the mixture brought to reaction temperature of 37 °C. 25 μL of 0.013 M Tris pH 6.0 containing l g/L BSA, 20g/L NaCl, l mL/L Tween 20, and 16.62 g/L NAD prewarmed to 37 °C is added to each of the wells. The mixtures are shaken for l minute then G6PDH activity is monitored by measuring the absorbance at 340 nm every 30 seconds for 3 minutes at 37 °C.

The dilution of conjugate giving an activity of between 200-250 mA/min is used for the EMIT assay.

6.14.1. Example: Conjugate of Peptide SEQ. ID. NO. 26 and G6PDH, designated EC1.

EC1 is diluted 1:10, 20, 40, 80, 160, 320 and 640 in a buffer of 0.218 M Tris pH 8.0 containing 1 g/L BSA and 7 g/L glucose-6-phosphate.

25 μL portions of diluted conjugate are added to appropriate wells of a microtiter plate. 100 μL of buffer (0.013 M Tris pH 6.0 containing l g/L BSA, 20g/L NaCl, 1 mL/L Tween 20) is added to each well and the mixture brought to reaction temperature of 37 °C. 25μL of 0.013M Tris pH 6.0 containing lg/L BSA, 20g/L NaCl, lmL/L Tween 20, and 16.62 g/L NAD prewarmed to 37 °C is added to each of the wells. The mixtures are shaken for 1 minute then G6PDH activity is monitored for 3 minutes by measuring the absorbance at 340 nm every 30 seconds for 3 minutes at 37 °C.

Results:

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Dilution EC1; 1:	Rate mA/min
10	800
20	393
40	218
80	123
160	63
320	35
640	18

A dilution of 1:40 was used for inhibition experiments with antibody and antigen (EMIT assay).

6.15. Determination of Amount of Antibody to Use

Commercially obtained antibody is diluted 1:10, 20, 40, 80, 160, 340 and 320 in a buffer of 0.013 M Tris pH 6.0 containing 1

g/L BSA, 20 g/L NaCl. 1 mL/L Tween 20. 100 µL portions of each dilution are added to wells of a microtiter plate. To each well is added 25 µL of diluted conjugate - dilution determined from activity experiments to give a rate of approx. 200-250 mA/min -- and the mixtures are incubated for 5-10 minutes at room temperature. The mixtures are then warmed to 37 °C and 25 µL of 0.013 M Tris pH 6.0 containing 1 g/L BSA. 20 g/L NaCl. 1 mL/L Tween 20, and 16.62 g/L NAD prewarmed to 37 °C are added to each of the wells. The mixtures are shaken for 1 minute, then G6PDH activity is monitored for by measuring the absorbance at 340 nm every 30 secs for 3 minutes at 37 °C.

The dilution of antibody giving an inhibition of 40-60% of G6PDH activity is used for the EMIT assay.

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6.15.1. <u>Example</u>

Goat anti-HBsAg is diluted 1:10, 20, 40, 80, 160, 340 and 320 in a buffer of 0.013 M Tris pH 6.0 containing 1 g/L BSA, 20 g/L NaCl, 1 mL/L Tween 20, 100 µL portions of each dilution are added to wells of a microtiter plate. To each well is added 25 µL of conjugate ECl diluted 1:40 and the mixtures are incubated for 5-10 minutes at room temperature. The mixtures are then warmed to 37 °C and 25 µL of 0.013 M Tris pH 6.0 containing 1 g/L BSA, 20 g/L NaCl, 1 mL/L Tween 20, and 16.62 g/L NAD prewarmed to 37 °C are added to each of the wells. The mixtures are shaken for 1 minute then G6PDH activity is monitored by measuring the absorbance at 340 nm every 30 secs for 3 minutes at 37 °C.

Results:

	Dilution of	Rate
	Ab: 1:	mA/min
5	10	125
	20	105
	40	135
	80	165
	160	200
10	320	217
	640	215
	no Ab	220

Dilution of goat anti-HBsAg to use for antigen experiments. 1:30.

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6.16. Dose Response to Antigen

Antibody dilution used is determined from the section, above. Conjugate dilution used is determined from previous experiments as used in the section, above.

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10-15 μ L of sample containing antigen is incubated for 10-15 minutes with 100ul of diluted antibody and 25 μ L of 0.013 M Tris pH 6.0 containing 1 g/L BSA, 20 g/L NaCl, 1 mL/L Tween 20, and 16.62 g/L NAD. The mixtures are then warmed to 37 °C and 25 μ L diluted conjugate pre-warmed to 37 °C is added. The mixtures are shaken for 1 minute then G6PDH activity is monitored for 3 minutes by measuring the absorbance at 340 nm every 30 secs for 3 minutes at 37 °C.

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The resulting dose response curve shows an increase in G6PDH activity (as a rate measurement; mA/min at 340 nm) proportional to the concentration of antigen added in the sample.

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6.16.1. Example

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10-15 μL of sample containing 0-200 ng/mL of rHBsAg antigen is incubated for 10-15 minutes with 100 μL of goat anti-HBsAg antibody (diluted 1:30) and 25 μL of 0.013 M Tris pH 6.0 containing 1 g/L BSA, 20 g/L NaCl, 1 mL/L Tween 20, and 16.62 g/L NAD. The mixtures are then warmed to 37 °C and 25 μL conjugate EC1 diluted 1:40 and pre-warmed to 37 °C are added. The mixtures are shaken for 1 minute, then G6PDH activity is monitored by measuring the absorbance at 340 nm every 30 secs for 3 minutes at 37 °C.

Results:

Conc. HBsAg ng/mL	Rate mA/min
0	100
3	112
6	122
12	140
2 5	165
50	185
100	205
200	220

An increasing rate of G6PDH enzyme activity is seen with increasing amounts of rHBsAg in the sample.

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6.17	Another Ex	cample of an EMIT Assay		
	6.17.1.	Conjugation of Binding	Peptides	10
		Enzyme Label		<u> </u>

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SEQ. ID. NO. 35 H₂N-LPGPPHLS-COOH

FW 816

FW

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SPDP

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The peptide of SEQ. ID. NO. 37 (31 mg) is prepared by air oxidation of the reduced form of the peptide in Kpi, pH 8 (reduction is effected by treatment with 1 mM tributyphosphine). Completion of oxidation is confirmed from a negative Ellman's test. Oxidized peptide is purified by reverse-phase HPLC, then lyophilized.

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An amount (1 mg) of the synthetic peptides, SEQ. ID. NO. 35 (ca. 1 umole) and SEQ. ID. NO. 37 (ca. 500 μ mole), is dissolved in 200 μ L PBS, 1 mM EDTA (PBSE - PBS containing 1 mM EDTA). The peptides are then thiolated using a 25-fold molar excess of a 100 mM (3.5 mg/250 μ L) solution of Traut's reagent in PBSE (50 μ L and 25 μ L, respectively). After allowing the resulting mixture to incubate in the dark for 1 hour at room temperature, the mixtures are exchanged into PBSE through a Sephadex G10 gel filtration column (Pharmacia, Piscataway, NJ).

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Separately, extensive dialysis of commercially obtained G6PDH enzyme in PBS provided approximately 1.2 mg G6PDH enzyme for dilution to a working concentration of 1 mg/mL (8 μ M, 390 U/mg). To 1.0 mL of this enzyme solution. G6P and NADH is added from fresh stocks (500 mM in PBSE) to provide a final concentration of 10 mM each. The enzyme is then allowed to react

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with 10-fold molar excess (80 µL) of a fresh 1 mM (0.5 mg/1.5 mL) solution of N-succinimidyl 3-(2-pyridyldithio)propionate (SPDP: Pierce, Rockford, IL) in carbitol. After a reaction period of 1h at room temperature, small molecules are removed by desalting on G10 in PBSE.

Labeled conjugates are formed by allowing appropriate amounts of the thiolated peptides to react with the activated enzyme in the refrigerator overnight.

Both preparations of conjugates are then harvested, filtered through a 1.0 um PFTE syringe filter, and brought to a volume of 3 mL by addition of an appropriate amount of Tris buffer. The resulting solutions of the labeled conjugates are then used in subsequent experiments as described further below.

6.17.2. <u>Preparation of Diluted Solutions of Labeled Conjugates</u>

Conjugates of peptides, SEQ, ID, NO, 37 and SEQ, ID, NO, 35 with G6PDH are designated EC4 and EC5, respectively.

Lyophilized samples of conjugates EC4 and EC5, as prepared above, are diluted 1:100, 200, 400, 800, 1600, 3200 and 6400 in a buffer of 0.218 M Tris (pH 8.0), containing 1 g/L BSA and 6 g/L glucose-6-phosphate.

Diluted solutions of each G6PDH-peptide conjugate (in 25 μ L portions) are added to appropriate wells of a microtiter plate. Then, 100 μ L of buffer (0.013 M Tris (pH 6.0), containing 1 g/L BSA, 20 g/L NaCl, 1 mL/L Tween 20) are added to each well. The resulting mixture is allowed to warm to the reaction temperature of 37 °C.

Next, 25 μL of 0.013 M Tris (pH 6.0), containing 1 g/L BSA,

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20 g/L NaCl. 1 mL/L Tween 20, and 16.62 g/L NAD (substrate for G6PDH) prewarmed to 37 °C are added to each of the wells. The mixtures are shaken for 1 minute, then G6PDH activity is monitored by measuring the absorbance at 340 nm every 30 seconds for 3 minutes at 37 °C.

The results are presented in the following Table.

Table 4. G6PDH Enzyme Activity

	RATE; mA/min at 340 nm												
		EC4		EC5									
DILUTION 1:	1	2	MEAN	1	2	MEAN							
BLANK	0	0	О	0	1	0							
100	221	214	217	333	312	322							
200	120	117	118	160	170	165							
400	65	64	64	9 0	91	90							
800	34	34	34	48	48	48							
1 60 0	17	18	17	25	31	28							
3200	10	13	1]	13	14	13							
640 0	5	5	5	7	7	7							

Consequently, to provide a signal level convenient for detection, a dilution of 1:150 is chosen for conjugate EC4, and a dilution of 1:200 is chosen for conjugate EC5.

6.17.3. <u>Inhibition of Labeled Conjugates by Affinity Receptor</u> Commercially obtained mouse anti-HBsAg is diluted 2500,

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1250, 625, 312, 160, 80 and 40 μ g/mL in a buffer of 0.013 M Tris (pH 6.0), containing 1 g/L BSA, 20 g/L NaCl, and 1 mL/L Tween 20. A portion (100 μ L) of each dilution is added to wells of a microtiter plate. Next, 25 μ L of a solution of either conjugate EC4 (1:150 diluted solution) or EC5 (1:200 diluted solution) are added to each well, and the resulting mixtures allowed to incubate for 1 hour at room temperature. Control wells using solutions of non-immune mouse IgG are also prepared for each conjugate.

The mixtures are then warmed to 37 °C and 25 µL of 0.013 M Tris (pH 6.0), containing 1 g/L BSA, 20 g/L NaCl, 1 mL/L Tween 20, and 16.62 g/L NAD prewarmed to 37 °C are added to each of the wells. The mixtures are shaken for 1 minute, then G6PDH activity is monitored by measuring the absorbance at 340 nm every 30 seconds for 3 minutes at 37 °C.

The results, presented in the Table, below, idicate that approximately 25% inhibition can be observed with both conjugates at the indicated concentration of antibody. The labeled conjugates are subsequently used in an EMIT assay for the detection of hepatitis surface antigen in a sample.

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TABLE 5

INHIBITION OF G6PDH-PEPTIDE CONJUGATE ACTIVITY
WITH INCREASING MOUSE ANTI-HBsag/NON-IMMUNE IgG

	ļ	RATES: mA/min at 340 nm													
			EC4*	; 1:150					EC5	; 1:200					
CONC Ab		MOU ANTI-H		7	IgG NON-IMMUNE			MOUSE ANTI-HBAAg			lgG NON-IMMUNE				
	1	2	MEAN	1	2	MEAN	1	2	MEAN	ı	2	MEAN			
0 4 0	198	203	200	184	210	197	254	282	268	264	267	265			
10	200	196	198	181	200	190	251 255	278	264	269	256	263			
166 312	203 193	206	204 192	212	202	207	251	265	258	267	270	268			
625	180	189	184	196	197	189 195	252 248	257	255 251	272	252	262			
1250 2500	166	171 152	149	17t 200	196	187 196	220 191	239	229	266 274	249	257			
										•/-	233	263			

* The labeled conjugate and antibody were pre-incubated for 1 hour at room temperature.

6.18. Method for Detection of Analyte of Interest With Labeled Mimetope

The following Example is for a method of determining the presence or absence of an hepatitis analyte in a sample by an antibody-mediated fluorescence enhancement affinity assay, such as one described by Wei, A-P, et al., in *Anal. Chem.* (1994) 66:1500-1506.

Hepatitis mimetope peptide of SEQ. ID. NO. 32 is allowed to react with tetramethylrhodamine-5-maleimide in 50 mM of phosphate buffer (pH 6) for 48 hours at 4 °C to make the labeled peptide. The

peptide is then reacted with 5-carboxyl-fluorescein succinimidyl ester in 50 mM borate buffer (pH 8.5) to double label the hepatitis peptide. FAB mass spectrometry is used to confirm the chemical identity of the doubly labeled peptide.

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The doubly labeled peptide is combined with a sample suspected of containing the hepatitis analyte, along with an appropriate amount of anti-hepatitis antibody.

The fluorescence activity of the sample is measured upon excitation with an ISS PC-1 fluorometer (ISS, Champaign, IL).

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The measured fluorescence enhancement is compared with the measured activity from a standard curve or the fluorescence observed from a control to determine the presence or absence of the hepatitis analyte in the sample.

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6.19 Method for Detection of Analyte of Interest with <u>Labeled Mimetope with Binding Profile for Antibody</u>

The following Example is for a method of determining the presence or absence of an hepatitis antibody in a sample by an affinity assay.

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Hepatitis mimetope peptide SEQ. ID. NO. 32, 35 or 37 is labeled with glucose-6-phosphate dehydrogenase as described above. FAB mass spectrometry is used to confirm the chemical identity of the labeled peptide. The labeled peptide is then used in an EMIT assay as described above to determine the presence or absence of hepatitis analyte in a given sample.

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The observed rates are compared with the measured activity from a control to determine the presence or absence of the antibody in the sample. Alternatively, the amount or concentration of analyte

can be determined quantitatively with the appropriate measurements and controls.

6.20. Kit in Accordance with the Invention

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Homogeneous immunoassay kit for performance of homogeneous assay of Example 18 comprises:

(a) a first container of functional surrogate. SEQ. ID. NO. 32, labeled with tetramethylrhodamine-5-maleimide and 5-carboxyl-fluorescein succinimidyl ester. The labeled peptide is capable of exhibiting an activity that is altered on binding of the labeled conjugate to the hepatitis affinity receptor and the activity can be measured and related to the amount of the analyte present in a given sample.

(b) a second container comprising goat or mouse anti-hepatitis surface antigen antibody.

6.21. <u>EMIT Kit</u>

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For an EMIT assay, a kit is provided including a first container of a functional surrogate (e.g., SEQ. ID. NOS. 35 or 37) labeled with G6PDII. A second container is also provided containing an antibody against the particular analyte of interest and, optionally, the G6P and NAD substrates for the GSPDH. If desired, these substrates can be present in a separate container.

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6.22. Recombinant DNA Construct Comprising a DNA Sequence Encoding a Functional Surrogate

A recombinant DNA construct is prepared which includes a DNA sequence encoding a hepatitis epitope or mimetope as obtained

by the methods of the invention This functional surrogate is capable of competing effectively for anti-hepatitis antigen antibody in the presence of the antigen. The recombinant DNA construct is made in accordance with methods known in the art such as those set forth in Sambrook et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Press (1989) using DNA inserts comprising the sequences set forth in this disclosure. A transforming vector, including the above DNA construct, a

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bacteriophage transformed by the vector, and a microorganism such as E. coli or yeast transformed by the vector or infected with the bacteriophage are made by methods known in the art such as those set forth in Sambrook et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Press (1989) or other references available to the skilled artisan, such as some of the patents mentioned herein. Preferably, such transforming vectors will include an origin of

replication functional in the host to allow for autonomous replication of the vector. Alternatively, the vector may integrate into the host

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In sum, the above disclosure teaches how to obtain, make and use functional surrogates having many uses, including, especially, use in homogenous enzyme immunoassays. The assay of the invention can advantageously detect macromolecular analytes, which have previously been difficult to assay for a number of reasons, as

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chromosome.

discussed above.

REFERENCES

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The purpose of the above description and examples is to illustrate some embodiments of the present invention without implying any limitation. It will be apparent to those of skill in the art that various modifications and variations may be made to the composition and method of the present invention without departing from the spirit or scope of the invention. All patents and publications cited herein are incorporated by reference in their entirety.

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MONOCLONAL ANTI-HBSAG BINDER SEQUENCES FROM R26 LIBRARY

RELEVANT	AMINO ACID SEQUENCES SEC	D. ID. NO.
FC23	MSMRSTVNVERRPAVAEPPAHLRINWGSR	176
FC41	VPTYWPSASILRSAETNGLHKLSHPLYSR	177
B26	ISSGLPSRLGCVSADAQTCHYHPIYNRSR	. 178
FC22	ACEIDPFYHPIYSAADQGARSDECIFPSR	179
FC21	DGSWWDMDLCSLPADCDALRSREKSRISR	180
FC32	LPGPPHLSVRHIPAESQNPTVDEAPAHSR	181
A28	TESAQRASSSTAASTHAVYGPPPNLSR	182
B13	ICAGASAGHQCRPAGPRHLDPSHSNGQSR	183
ClO	VQSVSSVGLMPYAAVSVHNNVSDHPLYSR	184
C24	VSA GTPTHTASLAAVNNYRHHPIYNPTSR	185
D20	FRPMQESLKAVDAAAAPPPYQFPMDDQSR	186
D7	HDLWCTGPRHLCPADMFPGTSNPSPPSSR	187
FC11	DAM SGGTGTSLDAAVIGPGHLFEYVDVSR	188
FC15	NFHAPFNHGEVETAASYLTDVPPHLLWSR	189
FC16	MAYFSSIGPVEHPAAGPGPLPRDFPPSSR	190

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TABLE 6 (con'd.)

MONOCLONAL ANTI-HBSAG BINDER SEQUENCES FROM R26 LIBRARY

SEOU	JENCES ALIGNED BY APPARENT MOTIFS	550	TD	
SEQUENT SEQUEN	ISSGLPSRLGCVSADAOTCHY ACEIDPFY VSAGTPTHTASLAAVNNYRH LP MAYFSSIGPVEHPAA DAMSGGTGTSLDAAVI HDLWCT ICAGASAGHOCRPA NFHAPFNHGEVETAASVITD MSMRSTVNVERRPAVAE TESAORA SSCTIVER	HPLY HPLY HPLY HPIY GPPHL GPRHL GPRHL VPPHL VPPHL ARSDE LESRE LESAE	PRDFPPSSR FEYVDVSR CPADMFPSTSNPSPPSSR DPSHSNGOSR LWSR RINWGSR SR CIFPSR KSRISR TNGLHKLSHPLYSR	184 177 178 179 185 181 190 188 187 183 176 182 179
		С а	pparent motif	198

TABLE 6 (con'd)

MONOCLONAL ANTI-HBSAG BINDER SEQUENCES FROM R2 ϵ LIBRARY

NUCLE	TIDE	SEQU	ENCES							SEC.	ID.	NO.
FC.23	CAC H	TCC S	TCG S	AGG P.	CTC L	CCC P	GGG G	CCC P	CCC P			191
	CAT H	CTG L	TCT	GTC V	CGG R	CAT H	ATT I	UCC P	GCG A			
	GAG E	AGT S	CAG Q	AAC N	CCC P	ACT T	GTT V	GAC D	GAG E			
	GCT A	CCC P	GCT A	CAT H	TCT S	AGA R						
FC.41	CAC H	TCC S	TCG S	AGT S	GTC V	CCT	ACT T	TAT Y	TGG W			192
	CCT P	AGC S	GCT A	TCT S	ATC I	CTC L	AGA R	TCC	GCG A			
	GAG E	ACC T	AAC N	GGG G	TTG L	CAC H	AAG K	CTT L	GAC D			
	CAC H	CCC P	CTT L	TAT Y	TCT S	AGA R			•			
B.26	CAC H	TCC S	TCG S	AGG R	ATT I	TCT S	TCT S	GGT G	TTG L			193
	CCT	TCG S	AGG R	CTG L	G GT G	TGC C	STG V	TCC	GCG A			
	GAC D	GCG A	CAG Q	ACC T	TGC	CAT H	TAC Y	CAC H	CCT			
	ATC I	TAT Y	AAC N	AGG R	TCT S	AGA R						
FC.22	CAC H	TCC S	TCG S	AGG R	GCT A	TGC C	GAG E	ATC I	GAT D			194
	CCT	TTT F	TAT Y	CAC H	CCT	ATC I	TAC Y	TCC S	GCG A			
	GCT A	GAC D	CAG Q	GGG G	GCT A	CGC R	AGT S	GAC D	GAG E			
	TGT C	ATT I	TTC F	CC3 P	TCT S	AGA R						

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TABLE 6 (con'd)

MONOCLONAL ANTI-HBSAG BINDER SEQUENCES FROM R26 LIBRARY

CAC H GAT D GAC	TC: S ATC M	2	G AG	C GA1	r GGG				SEO.	ID. NO
CAC H GAT D	S ATO M	S G GA	3		r GGG				<u>559.</u>	NC
H GAT D GAC	S ATO M	S G GA	3		r GGG					
D GAC	М		r ~~		G	AG: S	TG:	G TGG W		19
-	• тс•		r cro	TGT C	TCG S	CTC L	G CC			
	. 10.	GAT D	GCC A	TTG	CGC R	TCG S	CGC R	GAG E		
AAG K	AGC S	CGG R	ATC I	TCT S	AGA R					
H	TCC	TCG S	AGG R	CTC L	CCC P	GGG G	CCC	CCC		196
CAT H	CTG L	TCT S	GTC V	CGG R	CAT H	ATT	CCC P	GCG A		
GAG E	AGT S	CAG Q	AAC N	CCC P	ACT T	GTT V	GAC D	GAG		
GCT A	CCC P	GCT A	CAT H	TCT S	AGA R					
CAC H	TCC S	TCG S	ACA T	ACA T	GAG E	TCT	GCG A	CAG		197
AGA R	GCC A	TCT	TCA S	TCA S	ACC T	GCG A	GCC	TCC		
ACC T	CAC H	GCC A	GTC V	TAC Y	GGC G	CCT	CCC	CCT		
AAT N	CTT L	TCT S	AGA R					•		
CAC H	TCC S	TCG S	AGC S	ATT I	TGC C	GCT A	GGT G	GCT A		198
T CT S	GCT A	GGC G	CAC H	CAG Q	TGC C	CGT R	CCC	GCG		
GGT G	CCC P	GCG R	CAC H	TTG L	GAT C	CCG	AGT	CAC		
TCG S	AAC N	GGC G	CAG Q	TCT S	AGA R	-	<u>.</u>	п		
	CAC H GAG E GCT AAT AAT CAC TCG GG GCT TCG	CAC TCC H S CAT CTG H CTG CAC TCC A AGA AGT CAC TCC A AGA GCC A CAC TCC A TCC	CAC TCC TCG H S S CAT CTG TCT H L CAG E CCC GCT A CCC GCT CCC GCT CCC GCT CCC GCT CCC GCT CCC GCC CCC GCT CCC GCC CCC GCT CCC GCC CCC GCT CCC GCC	CAC TCC TCG AGG CAT CTG TCT GTC H L S S AGG GAG AGT CAG AAC E S Q AAC GAT CCC GCT CAT A P A H CAC TCC TCG ACA H S S S T AGA GCC TCT TCA AGA ACA ACC CAC GCC GTC T TCA ACC TCC TCG ACA ACC TCC TCG ACA ACC TCC TCG ACC T TCT TCA ACC TCC TCG ACC T TCT TCA ACC TCC TCG ACC T TCT TCT AGA ACC TCC TCG ACC T GCT GCC GCC CAC T GCT GCT GCC CAC GCG ACC GCG AAC GCC CAC CAC	CAC TCC TCG AGG CTC H S S R CTC CAT CTG TCT GTC CGG H L S V CGC H L S S AAC CCC H L S S AAC CCC GAG AAC CCC E S Q TCT TCA TCA A P A ACA T TCA AGA GCC TCT TCA TCA R A S S T TAC TAC T TCT TCT AGA ACC TCC TCG AGC ATT AAT CTT TCT AGA ACC TCC TCG AGC ATT TCT AGA ACC TCC TCG AGC ATT TCT AGA TCT TCT TCT TCA TCT TCT TCA TCA TCT	CAC TCC TCG AGG CTC CCC ACT TCG AGA ACA ACA ACA ACA ACA ACA ACA ACA A	CAC TCC TCG AGG CTC CCC GGG CAT ATT TCT AGA ACC ACT ATT TCT AGA ACC CCC ACT ATT TCT AGA ACC ACT ACT ACC ACT ACT ACC ACT ACT AC	CAC TCC TCG AGG CTC CCC GGG CCC H S S R L P GG CTC CCC ACT ATT CCC R H L S S R C TCT CCC ACT ATT CCC R H L S S Q N P T V D GAG E S Q N P T V D GAG E S S T T TCT AGA R ACA ACA ACA ACA ACA ACA ACA ACA ACA	CAC TCC TCG AGG CTC CCC GGG CCC CCC H S S R L P GG CAT ATT CCC GGG AA AC CCC ACT GTT GAC GAG E S Q N P A H S S S T T GC GCG CAT ATT CCC GAG AA P A T T T T T T T T T T T T T T T T	CAC TCC TCG AGG CTC CCC GGG CCC CCC H S S R L L P GG CAT ATT CCC GCG GAG AGC ATT ATT CCC GCG GAG AGC CAT ATT CCC GCG GAG AGC CAT ATT CCC GAG AGC AGC AGC AGC AGC AGC AGC AGC

TABLE 6 (con'd)

MONOCLONAL ANTI-HBSAG BINDER SEQUENCES FROM R26 LIBRARY

	ESTID	E SEQ	UENCE.	<u> </u>						SEQ.	ID. N	၁.
C.10	CAC H	TCC	TCG S	AGC S	GTT V	Č CAG	TCT S	GTG V	AGC S		1.	99
	AGC S	GTT V	G GG	TTG L	ATG M	CCT	TAC Y	GCC A	GCG A			
	GTG V	AGC S	GTT V	CAC H	AAC N	AAT N	GTC V	TCT S	GAC D			
	CAT H	CCG P	CTC L	TAT Y	TCT S	AGA R						
C.24	CAC H	TCC £	TC3 S	AGC S	GTG V	AGT S	GCG A	GGT G	ACC T		20) 0
	CCG P	ACC T	CAC H	ACG T	GCG A	AGC S	TTG L	GCC A	GCG A			
	GTG	AAT	AAC	TAT	CGT	CAC	CAT	CCC	ATT			
	v	N	И	Y	R	H	Н	P	I			
	TAT Y	AAC N	CCS P	ACT T	TCT S	AGA R						
D.20	CAC H	TCC S	TCG S	AGC S	TTT F	CGC R	SGG P	ATG M	CAG Q		20:	1
	gag E	AGT S	CTT	AAG E	GCC A	GTC V	GAC D	GCC A	GCG A			
	GCT A	GCG A	CCC P	CCC P	CCC P	TAC Y	CAG Q	TTC F	CCT P			
	ATG M	GAC D	GAT D	CAG ©	TCT S	AGA R						
D.7	CAC H	TCC S	TCG S	AGT S	CAC H	GAC D	TTG L	TGG W	TGT C		202	?
	ACT T	GGT G	CCG P	CGC F	CAT H	TTG L	TGC C	CCC	GCG A			
	GAT D	ATG M	TTC F	CCA P	GGC G	ACG T	AGC S	AAC N	CCC P			
	AGC S	CCG P	CCT	AGC S	TCT S	AGA R						

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TABLE 6 (con'd)

MONOCLONAL ANTI-HBSAG BINDER SEQUENCES FROM R26 LIBRARY

MUCI	JEOTIC	E SEQ	UENCE	<u>:s</u>						an-	
PC.1	.1									SEC.	ID. NO
	CA(S	TCC S	AG0	GAC D	GCC A	C ATS	TCC S	G GGT		20:
	GGT G	T	GGT G	ACC T	TCC S	CTA L	GAT D	GCC A			
	GTI V	Y ATT	GGT G	, CCC	GGC G	CAC H	CTT L	TTT			
	TAT Y	GTC V	GAC D	GTC V	TCT S	AGA R			~		
FC.15	5										
	CAC H	TCC S	TC3 S	AGC S	AAT N	TTT	CAC H	GCC A	CCT		204
	TTC F	AAC N	CAC H	GGT G	GAG E	GTC V	GAG E	ACC T	GCG A		
	GCC A	TCG S	TAC Y	TTG L	ACC T	GAT D	GTC V	CCC	CCC		
	CAT H	CTG L	CTC L	TGG W	TCT S	AGA R		∢ *	٤		
PC.16											
	CAC H	TCC S	TCG S	AGC S	ATG M	GCC A	TAC Y	TTT F	TCC		205
	TCC S	ATT I	GGT G	CCC	GTG V	GAG E	CAT	CCC	s GCG		
	GCT A	GGC G	CCC P	GGG G	CCC P	CTT L	CCC P	CGT R	A GAT		
	TTT	CCT	CCG P	TCC S	TCT S	AGA R	•	ĸ	D		

TABLE 7

MONOCLONAL ANTI-HBSAG BINDER SEQUENCES
FROM RBC LIBRARY

RELEVA	ANT AMINO ACID SEQUENCES	SEQ ID NO:
M10 M13 M20 M3 M4 M8 M12 M18 M23 M24 M29	CGGPEHLOVC CARGEVLPKC CSGPKHLOVC CGGRGASSRC COWWGGRDKC CDWKTVLPRC CSNGGPDHLC CDGPRHLSTC CEEGAVLPRC CKCHPLYGGC CEOGAVLAKC	206 207 208 209 210 211 212 213 214 215
SEQUEN: M24 M8 M13* M23 M29	CES ALIGNED BY APPARENT MOTIFS CKC HPLY GGC CDW KTVLPR C CAR GEVLPR C CEE GAVLPR C CEQ GAVLAK C	216 <u>SEQ ID NO:</u> 215 211 207 214 216
M20* M12 M18 M10*	CS GPKHL QVC CSNG GPDHL C CD GPRHL STC CG GPBHL QVC	208 212 213 206
M3 M4	CGGRGASSRC No apparent motif CQWWGGRDKC No apparent motif	209 210

internal war and the

TABLE 7 (con'd)

NUCLE	OTIDE S	EQUENCE:	<u> </u>						SEO	ID. NO.
M.10 CAC L CIT	TCC Q CAG	TCG V GTC	AGT C TGT	C TGT G GGA	G GGS S TCT	g ggg r aga	P CCG	E GAG	H CAT	217
M.13 CAC P	TCC E AAG	TCS C TGT	AGT G GGA	C TGT S TCT	A GCG R AGA	R AGG	G GGG	E GAG	v GTG	218 L TTG
M.20 CAC C CAG	TCC V GTG	TCG C TGT	AGT G GGA	C TGT S TCT	S AGT P. AGA	G GGG	p CCT	K AAG	H CAT	219 L TTG
M.3 CAC K AAG	TCC V GTC	TCG C TGT	agt gga	C TGT TGT	g GGG AGA	c G G C	R CGG	S AGC	I ATC	220 F TTC
M.4 CAC D GAT	TOC E AAG	TCG C TGT	AGT G GGA	C TGT S TCT	Q CAA R AGA	W TGG	w TGG	G GGG	g G G G	221 R CGG
M.8 CAC L CTG	TCC P CCG	TCG R AGG	AGT C TGT	C TGT C GGA	D GAT S TCT	W TGG R AGA	к AA G	‡ ACG	V GTT	222

TABLE 7 (con'd)

MUCI	ECTIDE	SEQUENCE	<u>:s</u>						SEQ.	ID NO.
M.12 CAC H CAT	TCC L CTC	TCG C TGT	AGT G GGA	C TOT S TOT	S TCT R AGA	n Taa	G GGG	G GGT	P CCG	223 D GAT
M.18 CAC S TCT	TCC T ACG	TCG C T GT	AGT G G GA	C TGT S TCT	D GAT R AGA	g ggg	P CCT	R CGT	H CAT	224 TTG
M. 23 CAC L TTG	TCC P C CG	TCG R CGG	AGT C TGT	C TGT G GGA	e gag s tct	E GAG R AGA	g GGT	A GCG	V GTG	225
M.24 CAC Y TAT	TCC G GGG	TCG G GGT	AGT C TGT	C TGT G GGA	K AAG S TCT	C TGT R AGA	h Cat	cai b	L CTG	226
M.29 CAC L TTG	TCC A 3CG	TCG K AAG	AGT C TGT	C TGT G GGA	E GAG S TCT	O CAG R AGA	g GGT	A GCG	V GTT	227

TABLE 8

POLYCLONAL ANTI-HBSAG BINDER SEQUENCES
FROM R26 LIBRARY

A3 D8 C2 A12	STS: SAD:	SEQUENCES SIGPLRHHAMTADSP	HTSIDEUCCE	SEO ID NO
D10 C14 B18 B13	LD: RC: STTI DC	WAEOGESTTSAGPR TREPSDGNCYAAAPLI SEDGLCYFGVDRGHWI ISKTKEREVTADPSI FSSAFFSGTARGPTI SSGPIKGALAADSOS	YTPDGLGMOA KAIPDLSDSR EPOLGVRNSL RSPHHPSAPT ATSSIONIGR	22 22 23 23 23. 23. 23.
A3 D8 C2 I	S ALIGNED B SAD WWWAEQJE XGFSSAFF LDF	Y APPARENT MOTI STSSIGPLR SNTPRGPLK STTSAGPRK STTSAGPRK STTTIATK SGTARGPIK SLTSSGPIK RCPSDGNCY RCPSDGLCY	HHAMTADSPHTGIDFHGGP YSADRLYTPDGLGMOA AIPDLSDSR DREVTADPSATSSIONIGE LGVLPSPOX GALAADSOSKPYSGPIMP AAAPLEPOLGVRNSL231 FGVDRGHWRSPHHPSAPT	SEQ ID NO. 228 229 230 233 234 235

TABLE 8 (con'd)

SELE	CIED NUC	LECTIDE	SEQUENC	<u>23</u>					SEQ.	ID. NO
B18	CAC A GCT P CCT P CCG	TCC F TTC T ACC Q CAG	TCG F TTT K AAG A GCC	AGT S AGT L TTG TCT	D GAC G GGT GGC R AGA	G GGG T ACG V GTG	F TTC A GCC L CTT	S AGC R CGC P CCC	S AGC G GGG S AGC	236
B13 CAC P CCG CAG P CAG P CCT	TCC I ATT S TCC S TCT	TCG K AAG K AAG R	AGT G GGG P CCT	S TCT A SCG Y TAC	L CTT L TTG S TCT	T ACG A GCC GC G	S TCT A GCG P CCT	S AGT D GAT I ATT	G GGC S TCT M ATG	237

TABLE 9

POLYCLONAL ANTI-HBSAG BINDER SEQUENCES FROM R&C LIBRARY

RELEVANT A	MINO ACID SEQUENCES	
P7 P8	CWLNWRGGTC CRGGDRHPGC	SEC. ID. NO.
P10 P19	CWEPYRGANC CGQICRQSLC	238 239 240
No apparent	t motifs from the above	241

TABLE 10 POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (A SERIES)

RELE	WANT AMINO ACID SEQUENCE	<u>S</u>	SEO I	D NO:
A1	HSSSSQHGGSAMFSLSSAAHSP			242
A3	HSSSFQLGSGGEALFKSAAALG:	PPGSRTPFI	ISR	243
A4	HSSSSASPTSVTFLROPAVSGG	RSLFQNLDI	PSR	244
A 6	HSSRDLFHGGQAMFNSAAVAAK:	SSGLISPDS	SSR	245
A7	HSSSKYGGMSLFQSQMTAGHHA	GTPPYTSRV	V SR	246
A8	HSSSSALFQSVAPLFSSAAPSNI	NDRSPKPFT	TSR	247
A9	HSSSLAYSPIGASLFOSAANNPS			248
All	HSSSLOLFTTALPWRDTAAPPMI			249
A12	HSSAAGGTSENONSWAAVAGG			250
A13	HSSSFRSSPHGRAMFQSAGNGS1			251
A14	HSSSTRTSQVSYGVSRPAAASH			252
A21	HSSSPWNVNAKNDDGMAAGRALI			253
A24	HSSRAFVPTFPMMTIRSAGRALI			254
A27	HSSTVSFKRPGFEQMAAGLQQ			255
SEOU	ENCES ALIGNED BY APPARENT	י אוריידידיפט	SEC ID NO.	
A24	HSSRAFVPTFPMMTIRS		ECRNDHASR	254
A13	HSSSFRSSP		SAGNGSFGNVPALSSSR	251
A6 A1	HSSRDLFH		SAAVAAKSSGLISPDSSR	244
A1 A3	HSSSSQH		LSSAAHSPAAHQATHTSSR	241
A3 A4	HSSSFQLGS HSSSSASPTSVTFLROPAVS	GGRSLFQ	SAAALGPPGSRTPFHSR	243
A7	HSSSKY		SOMTAGHHAGTPPYTSRWSR	244
A12	HSSSAAGGTSENONSWAAVA	GGASLFQ		246 250
A21	HSSPWNVNAKNDDGMA		OALNNGTSR	253
A.9	HSSSLAYSP		SAANNPSIPERTSDVSR	248
AB	нѕ		VAPLFSSAAPSNNDRSFKPFTSR	
A.11 F	HSSSLOLFTTALPWRDTAAPPML	SNSALFO		249
	HSSSTRTSQVSYGVSRPAAASHS	PORAFFO	VSF	252
A27 I	HSSSTVSFKRPGFEQMAAGLQQGQS	SINPTP	SR	255

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TABLE 10 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (A SERIES)

NUCL:	EOTIDE	SEQUE	NCES						SEO T	D NO:
A.1									220 1	
	CAC S AGC A GCT T ACG	TCC A GCC H CAT H CAC	TCG M ATG S AGC T ACC	AGT F TTT P CCC S AGC	S TCT S AGC A GCT S TCT	Q CAG L CTG A GCT R AGA	H CAC F TTC N CAT	G GGG S TCC Q CAG	G GGG A GCG A GCG	256
A.3										2==
	CAC G GGC A GCC T ACG	TCC G GGG A GCG P CCG	TCG E GAG L CTC F TTT	AGT A GCG G GGC H CAC	F TTT L CTT P CCC S TCT	Q CAG F TTT P CCG R AGA	TTG K AAG G GGG	G GGC S TCC S TCC	S TCC A GCG R CGC	257
A.4										
	CAC S AGC V GTG N AAC	TCC V GTC S AGT L CTC	TCG T ACT G GGG D GAT	AGC F TTT G GGG P CCC	S AGC L TTG R CGT S TCT	A GCG R CGG S AGT R AGA	S AGC Q CAG L CTC	P CCC P CCC F TTC	T ACC A GCG Q CAG	258
A.6										
	CAC G GGT V GTG S AGT	TCC Q CAG A GCT P CCC	TCG A GCT A GCT D GAC	R AGG M ATG K AAG S TCC	D GAT F TTT S AGC S TCT	L CTT N AAC S TCG R AGA	F TTC S TCG G GGT	H CAT A GCC L TTG	G GGG A GCG I ATC	259

TABLE 10 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (A SERIES)

NUC	CLECTI	DE SEQ	UENCES						SEO.	ID. NO.
A.7	•									<u> </u>
	CAC S AGT G GGC T ACG	L TTG H CAT	F TTT H CAT	Q	S TCG G GGG S	TATE Q CAC T ACC	F GG:	T GGT T ACC	A GCG Y	260
A.8	CAC S TCG A GCG K AAG	TCC V GTC P CCG P	TCG A GCC S TCG F TTC	S AGC P CCC N AAC T ACT	S AGT L CTG N AAT S TCT	A GCG F TTT D GAC R AGA	L TTG S TCG R CGG	F TTT S TCC S TCG	Q CAG A GCG P CCC	261
λ.9	CAC I ATC A GCT T ACT	TCC G GGT N AAC S TCC	TCG A GCT N AAC D GAT	S AGT S TCG P CCG V GTT	L TTG L TTG S AGC S TCT	A GCG F TTT I ATC R AGA	Y TAC Q CAG P CCC	S TCT S TCC R CGT	P CCC A GCG R CGT	262
A.11	CAC T ACT A GCG L CTT	TCC A GCT P CCG F	TCG L TTG P CCG Q CAG	AGC P CCG M ATG M ATG	L TTG W TGG L CTT S TCT	Q CAG R AGG S TCC R AGA	L CTG D GAC N AAC	F TTT T ACC S AGC	T ACC A GCG A GCC	263

109

TABLE 10 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (A SERIES)

NUCLE	OTIDE	SEQUE	1CES						SEO I	<u> </u>
A.12	CAC S TCG V GTT N AAT	TCC E GAG A GCC S AGC	TCG N AAT G GGG H CAC	AGC Q CAG G GGC R CGG	A GCT N AAC A GCG S TCT	A GCC S AGT S AGT R AGA	G GGG W TGG L CTT	G GGG A GCC F TTT	T ACC A GCG Q CAG	264
A.13	CAC H CAC G GGC A GCT	TCC G GGC N AAC L CTG	TCG R CGG G GGG S TCC	AGT A GCT S AGT S AGT	F TTT M ATG F TTT S TCT	R CGG F TTC G GGG R AGA	S TCC Q CAG N AAT	S TCG S TCC V GTC	P CCC A GCG P CCG	265
A.14	CAC V GTC A GCC F TTC	TCC S TCG A GCT F TTC	TCG Y TAT S TCG Q CAG	AGT G GGG H CAT V GTC	T ACG V GTC S TCG S TCT	R CGC S AGT P CCT R AGA	T ACT R CGT Q CAG	S TCC P CCC R AGG	Q CAG A GCG A GCT	266
A.21	CAC H GCC A GGG G AAT N	TCC S AAG K CGT R AAC N	TCG S AAC N GCC A GGG G	AGC S GAC D CTT L ACT T	CCT P GAC D TTT F TCT S	TGG W GGT G AAG K AGA R	AAT N ATG M CAG Q	GTG V GCC A GCG A	AAT N GCG A CTC L	267

FN: + Min was a war to war.

TABLE 10 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (A SERIES)

NUCL	EOTIDE	SEQUE	ENCES				,		SEO :	ID NO:
A.24	CAC R AGG D GAC T ACG	TCC T ACT R CGT D GAT	TCG N AAT S TCC L TTG	AGT L TTG F TTT D GAT	M ATG Q CAG T ACC S TCT	F TTT L AAG P CCT R AGA	Q TAG D GAC G GGT	E GAG T ACC Y TAT	H CAT A GCG R CGC	268
A.27	CAC R CGG G GGT N AAC	TCC P CCC L CTG P CCC	TCG G GGG Q CAG T ACC	AGT F TTT Q CAG P CCC	T ACG E GAG G GGA S TCT	V GTT Q CAG Q CAG R AGA	S AGC M ATG S AGT	F TTT A GCC S TCC	K AAG A GCG I ATC	269

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TABLE 11 POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (B SERIES)

REL	EVANT AMINO ACID SEQUENCES	SEQ. ID. NO.
B1 B10 B11 B13 B17 B18 B20 B22 B23 B25 B27	HS LELSINGTPVMGYTPRNIQEPKLSTDRNAR HSSSFSITSMGWSGATSAVSGGSSFWQHY/HSR HSSSCYFCDTGVGAPASAGTWSANGNNIHLTSR HSSSKDSFFQIDRLRSTAVNRIASNHPPMPNSR HSSSIDGIQGHSGLFGTAASRGIGNTVMFQASR HSSSGYKLHAGERNLAAAYAGTSSGERGLTSR HSSRQITAHPLTSVANLRGGDALFTQMRLHHSR HSSSLGNYNRGGMALFTAASSSRGQATERPVSR HSSSSMFCGAMFCQSSSAEHSRTTFKEANYLSR HSSSIVKQSVDVNLQVSADSPGTPASAFFQISR HSSSLFQENKLRGFLMSAGPST NRASTIDGSR HSSSASNGSSLFNDLKPAGGKLKLAPRATGISR	270 271 272 273 274 275 276 277 278 279 280 281
SEOE B1 B10 B17 B20 B22 B23 B25 B27 B11 B16	HS LEL HSSSFSITSMGWSAGTSAVS HSSSIDGIQGHSGLFGTAASRGI HSSRQITAHPLITSVANLR HSSSLGNYNE HSSSSMF HSSSIVKOSVDVNLOVSADSPGT HSSSAS HSSSCYFCDTGVGAPASAGTWSANGNNIHLTSE HSSSGGYKLHAGERNLAAAYAGTSSGERGLTSR HS LEL GGSFF VMGYTPKNIQEPKLSTDRNAR GGSFFW HYVHSR SKDSFFW HYVHSR SSSAFHSTAVNRIASNHPPHPHPNPNSE HISSSCH HSSSSKOFF HSSSSKAR HSSSSKAR H	SEC ID NO: 270 271 273 274 276 277 278

TABLE 11 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES
FROM R26 LIBRARY (B SERIES)

NUCLE	CTIDE S	BOUENCES	_					SEO	IF NO:
E.1	CAC T ACC N AAC D GAC	TCC P CCC I ATC R CGG	L CTA V GTG O CAG N AAC	E GAA M ATG E GAG A GCT	L S CTC AGC G Y GGA TAC P K CCC AAA R P CGA CCT	I ATA T ACG L CTC S TCG	N AAC P CCG S AGC R AGA	G GGA R CGG T ACA	282
B.IC	CAC M ATG V GTG H CAT	TCC S GGT S AGC Y TAT	TCG W TGG G GGT V GTG	AGC S TCC G GGT H CAC	F S TTC TCC G A GGT GCC S S TCG AGC S R TCT AGA	I ATT T ACC F TTC	T ACC S TCC W TGG	S TCC A GCG Q CAG	283
E.11	CAC T ACG G GGC I ATC	TCC G GGT T ACC H CAC	TCG V GTT W TGG L	AGT G GGC S TCT T ACG	C Y TGT TAT A P GCT CCT A N GCT AAC S R TCT AGA	F TTT A GCG G G	C TGT S TCC N AAC	D GAC A GCG N AAT	284
B.13	CAC CAG V GTG P	TCC I ATT N AAC M ATG	TCG D GAT R CGG P CCG	AGC R CGT I ATT N AAT	K D AAG GAT L R CTG AGG A S GCG TCT S R TCT AGA	S TCG S AGT N AAT	F TTC T ACC H CAT	F TTT A GCG P CCC	285

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TABLE 11 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (E SERIES)

NUCL	EOTIDE	SEQUENC	<u>ES</u>		DIBONE (E S	ERIES;			
B.17								SE	C ID NO:
	CAC G GGG T ACC T ACT	TCC H CAC A GCG V GTG	TCG S AGT A GCC M ATG	AGC GGT S TCT F	I D ATC GAT L F TTG TTT R G AGG GGT Q A CAG GCC	G GGT G GGG I ATT S TCT	I ATC G GGG R AGA	Q CAG N AAC	286
E.18	CAC H 70 CAT H GCT A CGT E	TCC S GCC A TAT Y GGT R	TCG S GGT G GCC A 10 GTI G	AGC S GAG E GGT G ACT T	GGT GGG G G CGG AAT R N ACC AGT T S TCT AGA S R	TAC Y TTG L TCC S	AAG K GCC A GST S	TTG L GCG A GAG E	287
E.20	CAC P CCT G GGA R CGC	TCC L CTA G SGA L CTG	TCT T ACG D GAT H CAC	AGA S AGC A GCC H CAT	Q I CAG ATT V A GTG GCT L F CTT TTC S R TCT AGA	T ACC N AAT T ACC	A GCA L CTC Q CAG	H CAC R CGC M ATG	288
	CAC H CGT R GCT A GAG E	TCC S GGT G AGC S CGG R	TCG S GGG G TCG S CCC P	AGT S ATG M TCT S GTT V	TTG GGT L G GCG TTG A L CGG GGT R G TCT AGA S R	AAT N TIT F CAG Q	TAT Y ACC T GCC A	AAT N GCG A ACG T	289

V.

TABLE 11 (com'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES
FROM R26 LIBRARY (B SERIES)

NUCLE	OTIDE S	EQUENCE	<u>s</u>						SEC	ID NO:
B.23	CAC A GCG E GAG A GCT	TCC M ATG H CAC N AAT	TCG F TTC S TCC Y TAC	AGC C TGT R CGT L CTG	S TCS CAG TAGC TCT	M ATG S AGC T ACG E AGA	F TTT S TCT F TTT	C TGC S TGC K AAG	G GGC A GCG E GAG	290
B.3	CAC V GTT D GAC F	TCC D GAT S AGC F TTC	TCG V GTT P CCT Q CAG	AGC N AAT G GGG I ATT	I ATC L TTG T ACG S TCT	V GTC Q CAG P CCG R AGA	K AAG V GTC A GCT	Q CAG S TCC S AGC	S TOT A GCG A GCC	291
B.25										292
	CAC H 70 AAG K GGT G ACG T	TCC S TTG L CCT P ATC I	TCG S AGG R AGT S 10 GGC G	AGT S GGC G ACC T GAT D	TTG L TTC F NAC TCT S	TTC F TTG L AAT N AGA R	CAG Q ATG M CGG R	GAG E TCC S GCG A	AAT N GCG A TCC S	
B.27	CAC H TO TCG S GGT G	TCC S CTT L GGG G ACG T	TCG S TIT F AAG K GGT G	AGC S AAT N CTT L ATT I	GCC A GAC D AAG K 10 TCT S	TCT S TTG L CTG L AGA	AAC N AAG K GCC A	GGG G CCC P	TCG S GCG A CGC R	293

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TABLE 12

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (C SERIES)

RELEVANT AMINO ACID SEQUENCES	SEO ID NO:
HSSSSLOTGGRAFFLTAGNPGGSTAIPGGLSR HSSRPLWOVDA KAADTAEYYLSHD SHPSDSR HSSRPSOSS FOLHSSTPAERMSTMRLNVPDASR HSSSLFOLRSSDKHPQAAGTSSASFGN SNHSR HSSSLFOLEAVPHWRRPADSHOLRAIHPHGDSR HSSSWRYSEVTAHDIPAGGAPLFQHERYLT SR HSSSWRYSEVTAHDIPAGGAPLFQHERYLT SR HSSSPGSL KSITDRNSAAAPAPSSNPLPSRSR HSSSLFONVVEGREMY AESPTTNILFHRHGSR HSPGGOVMWRLSNLDSADR KTKAHASGVSSR HSSSTMLCLNPLCWTAAGRLDTYTNPSTTSR HSPRPK SELDSVNYWPAGRAFFRDFFT LASR	294 295 296 297 298 299 300 301 302 303
SEQUENCES ALIGNED BY APPARENT MOTIFS C2 HSSSSLCT GGRAFFL TAGNPGGSTAIPGGLSR C29 HSPRPK SELDSVNYWP AERAFFR DFFT LASR C12 HSSR SQSS FQ LHSSTPAERMSTMRLNYPDASR C13 HSSSLFQ HSSSLFQ LRSSDKHPQAAGTSSASFGN SNHSR C18 HSSSLFQ LRSSDKHPQAAGTSSASFGN SNHSR C18 HSSSLFQ LRSDKHPQAAGTSSASFGN SNHSR C19 HCSSWRYSEVTAHDIFA AGAPLFQ EERYLT SR	SEQ ID NO: 294 304 296 301 297 298
C10 HSSEPLWOVDA KAADTAEYYLSHD SHPSDSR - no apparent motifs C11 HSSSPGSL KSITDRNSAAAPAPSSNPLPSRSR C26 HSPSGGOVMWRLSNLDSADR KTKAHASGVSSR C28 HSSSTMLCLNPLCWTAAGRLDTYTNPSTTSR	299 295 300 302 303

TABLE 12 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (C SERIES)

MITTO							44.1	OF SE	ries;		
NOC	<u>LEOTI</u>	DE SE	QUENC	ES						SEC	ID NO:
C.2	TC			CC TC	G AG	T TO	C AG		TC		305
	S AC			5 S	S	S	S		IG CAG L Ω		
	T	G				G TT F					
	GC A				G GG(G	G G	T TC	C AC			
	ATO I	2 CC P			CTC L	Ter	T AG	A CC	N TOG		
	AG# R	Α.						-	٦		
C.10	TCT	CA(TC(TCG S	AGG R	CCT P	CT3	TG(G CAG		306
	GTT V	GAT D	GCT A	NTG	AAG K	GCT A	GC3 A	GAC D			
	GCG A	GAG E	TAT Y	TAT Y	CTC L	TCT S	CAT H	GAC D	CGN		
	TCG S	CAC H	CCG P	TCG S	GAC D	TCT S	AGA R	CCA P	TCG S		
	AGA R							-	J		
C.12											
	TCT S	CAC H	TCC S	TCG S	AGG R	AGT S	CAG Q	AGT S	TCT S		307
	CNT	TTT F	CAG Q	TTG L	CAT H	TCC S	TCC S	ACG T	CCC		
	GCG A	GAG E	CGC R	ATG M	AGC S	ACC T	ATG M	CGC R	CTT		
	AAT N	GTG V	CCC P	GAT D	GCG A	TCT S	AGA R	CCN P	TCG S		
	AGA R										

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TABLE 12 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (C SERIES)

NUCLE	OTIDE	SEQUI	ENCES							SEO ID NO:
C.14	TCT S	CAC H	TCC S	TCG S	AGT S	CTT	TTC F	CAG Q	TTG I	308
	AGG E	TCT A	AGT V	GAC P	AAG H	CAC W	CCG R	CAG R	GCC P	
	GCG A	GGT D	ACC S	TCG H	TCC Q	GCG L	AGC R	TTT A	GGC I	
	AAT H	NCC P	AGT H	AAT G	CVC	TCT	AGA R	CCA P	TCG S	
C.18	AGA R									3 0 9
	TCT S	CAC H	TCC	TCG S	AGT S	CTT 1	TTT F	CAG Q	ATT	30.
	GAG E	GCG A	G TG V	CCT P	CAC H	TGG W	AGG R	CGG R	CCC	
	GCG A	GAC D	AGC S	CAT H	CAG Q	CTC L	CGG R	GCG A	ATT I	
	CAC H	CCC	CAT H	GGG G	GAT D	TCT S	AGA R	CCN P	TCG S	
	AGA R									
C.20										310
	TCT S	CAC H	TCC S	TCG S	AGT S	CTG L	TTT	TAG U	AAT N	
	GTG V	GTC V	GAG E	GGT G	CGT R	GAG E	ATG M	TAT Y	NCC	
	GCG A	GAG E	AGC S	CCT P	ACC T	ACC T	AAC N	TTA	CTT L	
	TTC F	CAT H	CGT R	CAT H	GGG G	TCT S	AGA R	CCN P	TCG S	
	AGA R									

TABLE 12 (con'd)

POLYCLONAL ANTI-FERRITIN EINDER SEQUENCES FROM R26 LIBRARY (C SERIES)

17710							12(1)	CSER	(IES)		
<u>NUC.</u>	LECTI	DE SE	OUENC	<u> </u>						SEQ 3	D NO:
C.2	TC S										311
	NC	C AA K		I AT	DA T	T GA!	T AGO R	3 AA N			
	GC(A			C CC(GCT.	r ccc	TCC S	TC S			
	CC.		G CCI		AGC R		T AGA	. CZI			
	AG <i>I</i> R	à									
C.23	TCT S	CAC H	TCC S	TCG S	AGT S	CTG L	TTT	TAG Q	AAT N		312
	g t g V	GTC V	GAG E	GGT G	CGT R	GAG E	ATG M	TAT Y	NCC		
	GCG A	GAG E	AGC S	CCT	ACC T	ACC T	AAC N	ATT I	CTT L		
	TTC F	CAT H	CGT R	CAT H	G GG G	TCT S	AGA R	CCN P	TCG S		
	AGA R										
C.26	TCT S	CAC H	TCC S	CCG P	AGT S	GGG G	GGG G	CAG Ç	GTG V		313
	ATG M	TGG W	CGT R	CTG L	AGC S	AAT N	TTG L	GAT D	TCC S		
	GCG A	GAC D	CGT R	ИСС	AAG K	ACG T	AAG K	GCT A	CAC H		
	GCT A	AGC S	GGC G	GTT V	TCN	TCT S	AGA R	CCA P	TCG S		
	AGA R										

TABLE 12 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY C SERIES;

MUCL	EOTID	E SEQ	UENCES	5	. 1.20	LIBRA	ury C	SERI	ES;		
C.28										SEQ	ID NO:
	TCT S	CAC H	TCC	TCG S	AGT S	ACT T	ATG M	TTG L	TGC C		314
	TTG L	AAC N	CCC P	CTT L	TGC C	TGG W	ACC T	GCG A	GCT A		
	GGC G	AGA R	CTC L	GAT D	ACC T	TAC Y	DOA T	AAT N	CCC P		
	TOT S	ACC T	ACG T	TCT S	AGA R	CCA P	TCS S	AGA R			
C.29											
	TCT S	CAC H	TCC S	CCG P	AGG R	CC3 P	AAG K	GNT	TCT S		315
	GAG E	CTT L	GAT D	TCG S	GTT V	AAT N	TAC Y	TGG W	CCC		
	GCG A	GG3 G	CGG R	GCC A	TTC F	TTT F	CGC R	GAC D	TTC F		
	P	ACT T	AAN	TTG L	GCG A	TCT S	AGA R	CCA P	TCG S		
	aga R										

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TABLE 13

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (D SERIES)

REL	<u>EVANT AMINO ACID SEQU</u>	ENCES		SEO	ID NO
	HSPRLYGGEALFOLL SAD HSSSSYGGRALFA ONPAV HSSSFMDIRKSPVAGTSAE HSSSFVLGTAGGSNVLSAG HSSSVTASGGEVLFKKTAA HSSRNLDEVAVGVEEGRGN	AVH THTD PSLHRPPA DRSPSSSC Y TGHIPS DTVADFAD LALFQQGA FSSNRHPS AFFKKFST RTFTRSEP PHLTTNIT	NNPILSR RATVSE TKC SP RHTSE OPRKSE OPRKSE NGPDSE SNAPSE IINSR DSHRSE PPHOSE		316 318 318 320 321 323 324 325 327
D23 D21 D3 D12 D17 D7 D20 D25 D26	HPSSSGGFVVSYRA HSPRLY HSSSSY HSSSTG HSSSFVLGTAGGSNVLS	RGNAFFK GGEVLFK GGSAAFQ GGEALFQ GGRALFA RVSTLFQ AGLALFQ AGSSSFFQ SSSAFFQ	KFSTIINSR KTAAFSSNRHPSSNAPSR NLTQEHPNT SE LL SADDRSPSSSCTKC SR QNPAVY TGHIPS RHTSR VQRAADPSLHRPPARATVSR CGANGPDSR FYNNGESRTSADRTPTRSEPDSHRS VNGRSLSSAGPHLTTNITPPHOSR		ID NO: 324 323 316 319 320 318 325 325
		THTD N	VPILSR - no apparent motif	S	327 317 321

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 $(A_{ij},A_{ij}$

TABLE 13 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (D SERIES)

NUCLE	OTIDE S	POUENC	ES						SEO. ID.	.си_
D.3										328
TCT SGTT V TTT F ACG	CAC H TCG S CAG O GAN	CCC P TAT Y AAT AN TCT S	TCG S CGG R CTG L AGA R	AGT S GCC A ACT T CCN P	AGT S GGG G CAG CAG S	GGC GGC GAG GAG E AGA R	GGG G TCC S CAT H	TTC F GCG A CCC P	GTG V GCG A AAC N	
D.6										329
TOT S ACT T CAT H ATT I	CAC H GGG G NAG CTG L	TOO SOUG RACT TOT S	TCG S ANC CAT H AGA R	NGG AAC N ACC T CCA P	ACT T AAG K GAC L TCG S	TGG W CCG P NGG AGA R	TCG S GCC A AAC N	ATT I GCG A AAT N	GAC D GTC V CCC P	
D.7										330
TCT S ACG T CCT P ACC	CAC H CTT L AGT S GTC V	TOO STATE CTO TOT S	TCG S CAG Q CAC H AGA R	AGC S GTT V AGG R CCA P	ACT TAG COG FTCG S	GGC G AGG R CCG P AGA R	AGG R GCC A GCG A	GTT V GCG A CGC R	AGT S GAT D GCC A	
D.12										331
TCT S GCG A GAT D AAG K	CAC H CTT L GAT D TGT C	TCC S TTC F CGC R GAN	CCG P CAG Q TCT S TCT S	AGG R CTG L CCC P AGA R	TTG L CTT L AGC S CCA F	TAC Y NTG AGC S TCG S	GGC G TCC S AGT S AGA R	GGC G GCG A TGC C	GAG E ACG T	234

TABLE 13 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (D SERIES)

NUCLE	SEQ. II	<u> 110 .</u>								
D.17										332
TCT S AGG R GCG A TCG S AGA R	CAC H GCG A GTG V NTT	TCC S TTG L TAT Y CGT R	TCG S TTT F NCG CAC H	AGC S GCG A ACG T ACG T	TCG S ANC GGT G TCT S	TAC Y CAG Q CAC H AGA R	GGC G AAC N ATC I CCA	GG C G C C P C P C P C P C S		
D.19										333
CAC H AAG K GAG E CCC	TOC S TOC S GAC D CGG R	TCG S CCC P ACC T AAG K	AGC S GTT V GTT V TCT S	F GCC A GCT A AGA R	ATG M GGC G GAC D	GAT D TAT Y TTC F	ATC I TCC S GCG A	AGG R GCG A GAC D		
D.20						•				334
CAC H GGC G GCT A GAC D	TCC S GGT G CTC L TCT S	TCG S AGC S TTC F AGA R	AGT S AAT N CAG Q	TTT F GTG V CAG Q	GTG V TTG L GGC G	CTT L TCC S GCC A	GGG G GCG A AAT N	ACG T GGT G GGC G	GCC A CTC L CCT P	
D.21										335
CAC S GTG V AGC	TCC S TTG L TCC	TCG V TTT F AAC	AGC T AAG K CGG	GTT A AAG K CAT	ACC S ACC T CCG	GCC G GCG A AGC	AGC G GCG A TCT	GGC G TTT F AAC	SGC G SCT	GAG E CCG
S TCT S	s Aga R	N	R	Н	p	S	S	N	Ā	F

TABLE 13 (con'd)

POLYCLONAL ANTI-FERFITIN BINDER SEQUENCES FROM R26 LIBRARY (D SERIES)

NUCLE	OTIDE_	SEQUENC	<u>ies</u>						SEQ. ID.	<u>ND.</u>
D.23										336
CAC H GTA V TTT F TCT S	TCC S GGG G TTC F AGA P	TCG S GTG V AAG K	agg R GAG E AAG K	AAT N GAA E TTT F	TTG L GGC G AGC S	GAC D CGC R ACT T	GAA E GGC G ATC I	GTT V AAT N ATT I	GCA A GCC A AAT N	336
D.25										227
CAC H AAT N ACG T CGG R	TCC S GGG CCC P TCT S	TCG S GAG E ACG T AGA R	AGC S AGC S AGG R	TTT F CGG R TCC S	TTT F ACT T GAG E	CAG Q TCC S CCG P	TTT F GCG A GAT D	TAC Y GAT D AGT S	AAC N CGT R CAC H	337
D.26										2.0.0
CAC H AAC N CAT H CAG O	TCC S GGG G CTC L TCT S	TCG S AGG R ACC T AGA R	AGC S AGC S ACC T	TCG S CTG L AAC N	GCC A TCT S ATC I	TTC F TCC S ACC T	TTT F GCG A CCG P	CAG Q GGC G CCC	STC V CCG F CAC H	338
D.29										339
CAC H GAT D CCT P AGG R	TCC S GAT D ATC I TCT S	TCG S CGG R ACT T AGA R	AGG R GTG V AAT N	AAT N AAT N ATG M	TCT S GCA A TTC F	TTT F ACC T CAG Q	TTT F GCG A CAC H	TGG W GAG E TCC S	GGT G CCC P AAG K	<i>907</i>

TABLE 14 POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (AD SERIES)

RELEVANT AMINO A	CID SEQUENCES		SEQ ID NO:
AD3 HSSSKYGGMSL AD14 HSSSLFQSTPG	FCSOMTAGHHAGTPPY	TSRWSR	340
AD15 HSSSYNVVAGE	RVRLMPAANDGISSTP	GRIPSR	341
ADIS RESCROYMEN	AFFRDTAVNTAYPOTA	FETRSR	342
AD16 HSSSSSVTVVR AD17 SHSSRQITAHP	ANSAISAVRISNIA L	HTDRSR	343
AD18 HSSSPDSVGGH	LISVANLRGGDALFTQ	MRLHSR	344
AD22 HSSSMFQEHRTI	SFFRSSAGSHHRAHAR	APGNSR	345
AD26 HSSSIRTPFSRI	NACT NEEDS CAREADERS.	TDLHSR	346
AD27 HSSSGSSMFQVI	DEMICEDER VAPILL	P_STSR	347
AD29 HSSSSLFQRHNI	DK A A DEM T THE DAMPE A THE	RRYDSR	348
ime: Housest Okine	COMMERANTERUSA	indsk	349
SEQUENCE: ALIGNEI		<u>FIFS</u> R DTAVNTAYPQTAFETRSR	SEQ IT NO:
AD17 SHSSRQITAHPI		<u>C</u> DIAVNIAIPQTAFETRSR I QMRLHSR	342
		E QMRDASK ∑ SSAGSHHRAHARAPGNSR	344
AD3		2 SQMTAGHHAGTPPYTSRWSE	345
AD27	HSS SGSSMPC	VDRVVSSADIKMPPVHIRKYDSR	340
AD22	HSSSMFC	EHRTNLOKNTADRSFTPGYRTDLH.	348
AD29	H SSSSLFC	RHNRVDMMPAAHNPPKDSATLHGS	SR 346
AD14	HSSSLF		R 349 SR 341
AD16 HSSSSSVTVVFA AD26 HSSSIRTPFSRN	NSATSAVKTSNTA LH YELVSAGASVAPLLLP	TDRSR - no apparent motifo	3 343 347

TABLE 14 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES
FROM R26 LIBRARY (AD SERIES)

NUCLEOTIDE SEQUENCES											
AD.3									SEO I	I NO:	
TCT S	CAC H	TCC S	TCG S	AGC S	AAG K	TAT Y	GGT G	GGT G	82 ATG	350	
AGT S	TTG L	TTT F	CAG Q	TCG S	CAG Q	ATG M	ACC	GCG	M GGC		
CAT H	CAT H	GCG A	G G G G	ACC T	CCC P	CCG	TAT Y	A ACG	G TCC		
AGG R	TGG W	TCT S	AGA R	CCT P	TCG S	AGA R	i	T	Q)		
AD.15	5										
TCT S	CAC H	TCC S	TCG S	AGC Y	TAT N	AAC V	gtg V	GTT	GCT	351	
GGG R	CGC A	TTT F	TT ⊂ F	CGG R	GAC D	52 CCC P	GCG A	A GTC V	G TCC		
AAC T	ACC A	GCC Y	TAC P	O CCI	CAG T	ACT A	GCC F	TTC E	n Gag		
ACG R	CGG S	AGA R	CCT P	TCG S	AGA R		-		T		
A.16											
TCT S	CAC H	TCC	TCG	AGT	Tar	AGT	_		352		
g r g	n CGG	S	S	S	S	AG1 S 52	GTG V	ACG T	G T G V		
v	R	GCG A	AAC N	TCG S	GCT A	ACG T	TCC S	GCG A	GTG V		
AAG K	AAG T	ACC S	TCC N	AAC T	ACG A	GCG	NAG L	CTT H	CAT		
ACG D	GAC R	AGG S	AGA R	CCT P	TCG S	AGA R		п	Т		

TABLE 14 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES
FROM R26 LIBRARY (AD SERIES)

NUCLE	STIDE .	SEQUENC	EES						SEC :	II NC:
AD.18									3 5 3	3
TCT S	CAC H	TCC S	TCG S	AGT S	CCG P	GAT D 52	AGC S	GTC V	GGG G	
GGG G	CAT H	TCG S	TTT F	TTT F	AAG K	TCG S	TCC S	GCG A	GGC G	
TCT S	CAT H	CAC H	CGT R	GCG A	CAT H	GCG A	C G C R	GCG A	CCG P	
GGC G	AAT N	TCT S	AGA R	CCT P	TCG S	AGA R				
AD.26										354
TCT S	CAC H	TCC S	TCG S	AGT S	ATT I	AGG R	ACG T	CCT P	TTT F 52	
TCT S	CGG R	TAA N	TAC Y	GAG E	TTG L	GTT V	TCC S	GCG A	GGC G	
GCT A	AGC S	GTC V	GCT A	CCT P	CTC L	CTC L	T T G L	CCC P	ATC I	
TCC S	ACT T	TCT S	AGA R	CCT P	TCG S	AGA R				
AD.27										355
TCT S	CAC H E2	TCC S	TCC S	agt S	GGG G	AGT S	TCG S			
ATG M	TTC F	TAG Q	GTG V	GAT D	CGT R	GTC V	GTC V			
TCT S	TCC S	GCG A	GAT D	ATC K	AAG M	ATG P	CCC P			
CCC P	GTG V	CAC H	ATT I	CGC R	aag K	TAT Y	GAT D			
TCT S	AGA R	CCT P	TCG S	AGA R						

TABLE 14 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM R26 LIBRARY (AD SERIES)

MUCLE:	OTIDE .	SEQUENC	CES						CEO 75 350
AD.29									SEO II NO:
TCT S	CAC H	TCC S	TCG S	AGT S	TCC S	CTG L	TTT F	CAG Q	356
82 CGT R	CAC H	AAC N	AGG R	GTC V	GAT D	ATG M	ATG M	CCC P	
GCG A	GCT A	CAC H	AAC N	CCG P	CCG P	AAG K	GAT D	TCT	
GCC A	ACG T	CTC L	CAC H	G3G G	TCT S	AGA R	CCT	TCG S	
AGA								_	

TABLE 15

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM D38 LIBRARY

RELEVANT	AMING ACID SEQUENCES	
T15-4 T15-10 T15-17 T15-27 T15-29		357 358 359 360 361 362
T15-29 T15-10 H	ALIGNED BY APPARENT MOTIFS HSSSSAGNC ESRAGDSAA GRGGITGV GGGAMFO ESDMPGRIS RGRAMFK HSSSMFO EGKRRGLPGWICNEGHSHAIHNPNLNQCPDPSPGP	361 358 357 360 359
T15-30 H	SSRCLGSNTGEGRTWGTSSQINLDAIPNYTTPHIRQTVPYSSR - no apparent motif	362

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No Commence

Section 1985 Section 2015

e de la companya del companya de la companya del companya de la companya del la companya de la c

TABLE 15 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM D36 LIBRARY

FROM D36 LIBRARY NUCLEOTIDE SEQUENCES												
	5 - 4								SEO	ID NO:		
CAC H	TCC S	TCG S	AGG R	GGT G	GGG G	GGC G	ATC	ACC	121 GGG	363		
GTC V	G G	GGG G	GGT G	GCG A	ATG M	TTT F	I CAG Q	T TCC	G 91 CGT			
CCC P CGC	TCT S	GTT V	TTC F	AAC N	GCC A	ATT I	AGC S	S AAC	R 61 AAT			
R CAC	GGC G	CAC H	ACG T	ATT I	CCC P	GAC D	ACT T	N TTT F	N 31 CCC			
H	ACT T	TCT S	AGA R	ATC I	GAA E	GGT G	CGC R	GCT A	P 1 AGA			
T.15	-10							••	R			
CAC H GCT	TCC S 121	TCG S	AGA R	GCG A	GGG G	GAC D	AGT S			364		
A	GCG A	G G C G	GGC G 91	ATG M	GCG A	CTT	TTT					
CGC P	GAT D	GTC V	CCG P	CTG L	TCG S	ATT I	F CGT R					
GAC D	61 GCC A	AGG R	CCC P	CCT P	GCC A	CAC	CCT					
AAT N	AGC S	AGC S	CAT H	CTT L	ATC	H GAT D	P TGG					
AGC S	ACT T	TCT S	AGA R	ATC I	GAA E	G GT	W CGC					
GCT A	AGA R				-	J	R					

TABLE 15 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM D38 LIBRARY

NUCLE	ECTIDE	SEQUEN	CES						SEO	ID NO
T.15	-17									365
CAC H 121	TCC S	TCG S	AGC S	ATG M	TTC F	CAG Q	GAG E	GGT G		
AAG K	C G G R 91	AGG R	G GT G	TTG L	CCG P	GGT G	TGG W	ATC I		
TGC C	AAT N	GAG E 61	GGC G	CAT H	TCT S	CAC H	GCC A	ATC I		
CAC H	AAT N	CCC P	AAT N 31	CTC L	AAC N	CAG Q	TGT C	CCC P		
GAC D	CCG P	AGT S	C C G P	GGC G	CCT P	TCT S	AGA R	ATC I		
GAA E	GGT G	CGC R	GCT A	AGA R						
T.15-	27									366
CAC H	TCC S	TCG S	AGT S	GAC D	ATG M	CCG P	GGG G	CGG R 79	ATT I	TCT S
CGG R	G GT G	CGC R	GCC A	ATG M	TTC F	AAG K	GAG E	GTT V	CAC H	GCC A
ACT T CCT P	ACC T CAT H	CAT H CAT H	GCC A ACC T	GAT D CCG	GAG E TCT	GTG V AGA	GGC G	GGC G	ACG T	AAC N

TABLE 15 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM D38 LIBRARY

NUCLECTIDE SEQUENCES SEC IT NO									D_NO:	
T.15-	29									367
CAC H	TCC S	TCG S	AGT S	TCG S	GCG A	GGC G	AAC N	TGT C	121 TGC C	
CGG R	GGT G	AGC S	CTT L	TTC F	TGC C	TCT S	TGC C	GGT G	91 GAG E	
CGT R	ACT T	G GT G	ATG M	GAC D	GCC A	ATC I	ACC T	CCC P	61 CAT H	
CCG P	CAT H	ATC I	CTC L	CAC H	CGC R	GGG	AGC S	TCC S	31 TCT S	
GCC A	GCC A	TCT S	AGA R	ATC I	gaa E	GGT G	CGC R	gct A	AGA R	
T.15-	30			∢						368
CAC H	TCC S	TCG S	AGG R	CAG Q	CTG L	G GT G	TCG S	AAT N	121 ACG T	
GG G G	GAG E	G GT G	CGG R	ACT T	TGG W	GGT G	ACT T	TCC S	91 TCG S	
CAG Q	ATC I	AAC N	CTG L	GAC D	GCC A	ATC I	CCT	AAC N	61 TAC Y	
ACC T	ACC T	P CCC	CAC H	ATT I	C G G R	CAG Q	ACG T	GTT V	31 CCG p	
TAC Y	TCC S	TCT S	AGA R	ATC I	GAA E	GGT G	CGC R	GCT A	i AGA R	

TABLE 16

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM DC43 LIBRARY

AMING A	ACID SEQUENCES	SEO ID	NO.
T20-1 T20-5 T20-13	HSSSNYGGADRAWGGRSLFTSAVTGCGNSPRNDRDERRPNTRTSNVTSP HSSRTAKEGCSGGASLFLELRAQCGCGAHRNTPPSHCLPVETKNCDDSP HSSSINDSGSRTWSGGCGISRDGARALFLDDPSRDPLSR	<u> </u>	36 37 37
SEQUENC	ES ALIGNED BY APPARENT MOTIFS	SEO ID	110
T20-5 F	HSSSNYGGADRAW GGRSLFT SAVTGCGNSPRNDRDERRPNTRTSN HSSRTAKEGCS GGASLFL ELRAQCGCGAHRNTPPSHCLPVETH HSSSINDSGSETWSGGCGISRD GARALFL DDPSRDPLSR	SEO ID WTSR INCDDSR	3.00

TABLE 16 (con'd)

POLYCLONAL ANTI-FERRITIN BINDER SEQUENCES FROM DC43 LIBRARY

NUC	NUCLEOTIDE SEQUENCES SEQ ID No:														
T.2	0-1												-		372
CAC H	TCC S	TCG S	AGC S	AAT N	TAC Y	GGT G	GGC G	GCG A	GAT D						J / Z
A GG R	GCG A	TGG W	GGT G	GGG G	CGG R	TCG S	CTG L	TTC F	ACG T						
AGC S	GCT A	GTG V	ACC T	GGT G	TGT C	GGT G	AAC N	TCC S	CCC P						
CGT R	AAC N	GAT D	AGG R	GAC D	GAG E	CGC R	CGT R	CCT F	AAC N						
ACG T	AGG R	ACT T	AGT S	AAT N	GTT V	ACC T	TCT	AGA R	ATC I						
GAA E	GGT G	CGC R	GCT A												
T.2	0 - 5														
CAC H	TCC S	TCG S	AGA R	ACG T	GCT A	AAG K	GAG E	GG3 G	AGT S	GTG V				•	373
GGC G	GGG G	GCC A	AGC S	CTG L	TTT F	TTG L	GAG E	CTT L	AGG P.	GCC A					•
CA G Q	TGT C	GGT G	TGT C	GGT G	GCT A	CAC H	CGT E	CAA N	ACC T	CCG P					
CCG P	TCG S	CAC H	TGC C	TTG L	CCT P	GTT V	GAG E	ACA T	AAG K	AAT N					
TGT S	GAT D	GAC D	TCT S	AGA R	ATC I	GAA E	GGT G	CGC R	GCT A	AGA R					
T.20	0-13													-	374
CAC H	TCC S	TCG S	AGT S	TTA	AAT N	GAC D	AGT S	GGT G	AGC S	AGG R				-	, , •
ACG T	TGG W	TCG S	GGT G	GGT G	TGT C	GGT G	ATC I	TCC S							
CGG R	GAT D	GGC G	GCC A	CGC R	CGC R	GCC A	CTT L	TTC F	CTG L	GAC D					
GAT D	CCC P	TGC C	CGC R	GAC D	CCT P	TTG L	TCT S	AGA R							
ATC I	GAA E	GGT G	CTC R	GCT A	AGA P.										

WHAT IS CLAIMED IS:

1. A method of determining the presence or absence of an analyte of interest in a sample by an affinity assay comprising:

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(a) providing (i) a labeled conjugate comprising at least one label attached to a functional surrogate capable of competing effectively with an analyte of interest for a limiting amount of an affinity receptor for said analyte, said labeled conjugate exhibiting an activity that is altered on interaction of said labeled conjugate with said affinity receptor and which activity can be measured and related to the presence or absence of said analyte in a given sample, and (ii) said affinity receptor;

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(b) combining said labeled conjugate and affinity receptor with a sample suspected of containing said analyte to provide a measure of said activity;

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(c) measuring said activity; and

(d) relating said activity to the presence or absence of said analyte in said sample.

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2. The method of claim 1 in which said interaction is a binding interaction.

3. The method of claim 1 in which said functional surrogate is further characterized as exhibiting a competitive binding profile that is substantially similar to that exhibited by said analyte for said affinity receptor.

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4. The method of claim 1 in which said functional surrogate is further characterized as exhibiting a selective binding affinity (K_a) for said affinity receptor which is substantially similar to that exhibited by said analyte.

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- 5. The method of claim 1 in which said relating step provides a quantitative measure.
- 6. The method of claim 1 in which said functional surrogate is obtained by screening a random peptide library with an affinity receptor of said analyte.
 - 7. The method of claim 6 in which said random peptide library comprises a plurality of peptides whose structures are not dictated by the primary sequence of said analyte.

8. The method of claim 1 in which the molecular structure of said functional surrogate corresponds to an epitope of said analyte.

- 9. The method of claim 8 in which the structure of said epitope was previously unknown.
- 10. The method of claim I in which the molecular structure of said functional surrogate differs from that of a known epitope of said analyte.

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11. The method of claim 10 in which said molecular structure does not include a primary sequence of eight or more consecutive amino acid residues which can be found along the naturally occurring sequence of said analyte.

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- 12. The method of claim 1 in which said functional surrogate has a molecular weight of 2000 daltons or less.
- 13. The method of claim 1 in which said functional surrogate comprises a peptide.

14.	The method of claim 1 in which said analyte is a
hapten.	
15.	The method of claim 1 in which said analyte is an
antigen.	
16.	The method of alain 1:
10.	The method of claim 1 in which said analyte is an
antibody.	

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- 17. The method of claim 1 in which said combining step comprises forming an affinity receptor-labeled conjugate complex.
- 18. The method of claim 17 in which said combining step further comprises displacing said labeled conjugate from said complex with said analyte.
- 19. The method of claim 1 in which said combining step comprises providing competition among said analyte and said labeled conjugate for said affinity receptor.

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20. The method of claim 1 in which said combining step comprises forming an affinity receptor-analyte complex.

- 21. The method of claim 20 in which said combining step further comprises forming an affinity receptor-labeled conjugate complex.
- 22. The method of claim 1 in which said sample is obtained from a biological fluid selected from the group consisting of urine, semen, saliva, sweat, blood, serum, plasma, cerebrospinal fluid, tears, vaginal or nasal fluids.
 - 23. The method of claim 1 in which said sample is obtained from a cell-free extract.

24. The method of claim 1 in which said label is selected from the group consisting of a chromogenic agent, UV absorber, fluorescent molecule, a chemiluminescent compound, an enzyme, an enzyme fragment, an enzyme substrate or a group having the potential for exhibiting at least one of the above-recited activities.

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25. The method of claim 24 in which said label comprises an enzyme.

- 26. The method of claim 25 in which said enzyme exhibits glucose-6-phosphate dehydrogenase (G6PDH) activity.
- 27. The method of claim 1 in which said combining step comprises (i) mixing said affinity receptor and sample, and (ii) adding said labeled conjugate to the resulting mixture.

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- 28. The method of claim 1 in which said activity is measured as a rate of change.
- 29. The method of claim 1 in which said analyte is a polypeptide, a polysaccharide, a polynucleotide, a glycoprotein or a lipid-containing macromolecule.
- 30. The method of claim 1 in which said analyte is a fertility/pregnancy-related hormone, is related to an infectious disease, is a cardiac marker or a tumor marker.

31. The method of claim 1 in which said analyte is associated with a bacterial or viral infectious agent.

The method of claim 1 in which said analyte is ferritin. follicle stimulating hormone (FSH), thyroid-stimulating hormone (TSH), human growth hormone (hGH), immunoglobulin E (IgE), prolactin, parathyroid hormone (PTH), human placental lactogen (HPL), human chorionic gonadotropin (hCG), human leutinizing hormone (hLH), cytomegalovirus (CMV), chlamydia, streptococcus A, rubella, toxoplasma, herpes, hepatitis, CK-MB, myoglobin, myosin light chain, troponin, carcinoembryonic antigen (CEA), alpha-fetoprotein (AFP), prostrate specific antigen (PSA), CA125 (a tumor marker).

33. The method of claim 1 in which said analyte is an allergen.

34. The method of claim 1 in which said analyte has a molecular weight in excess of about 100,000 daltons.

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- 35. A method of determining the presence or absence of an analyte of interest in a sample by an homogeneous enzyme affinity assay comprising:
- enzyme attached to at least one functional surrogate of an analyte of interest, said functional surrogate capable of competing effectively with said analyte for a limiting amount of an affinity receptor for said analyte, said enzyme conjugate exhibiting an activity that is altered on interaction of said enzyme conjugate to said affinity receptor and which activity can be measured and related to the amount of said analyte present in a given sample; (ii) said affinity receptor, and (iii) a substrate for said enzyme:
- (b) combining said enzyme conjugate, affinity receptor, and enzyme substrate with a sample suspected of containing said analyte to provide a measure of said enzyme activity;
 - (c) measuring said enzyme activity; and
- (d) relating said enzyme activity to the presence or absence of said analyte in said sample.

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- 36. A method of determining the presence or absence of an analyte of interest in a sample by an homogeneous fluorescence polarization affinity assay comprising:
- (a) providing (i) a labeled conjugate comprising a fluorescent material attached to a functional surrogate of an analyte of interest, said functional surrogate capable of competing effectively with an analyte of interest for a limiting amount of an affinity receptor for said analyte, said labeled conjugate exhibiting an activity that is altered on interaction of said labeled conjugate with said affinity receptor and which activity can be measured and related to the presence or absence of said analyte in a given sample, and (ii) said affinity receptor;
- (b) combining said labeled conjugate and affinity receptor with a sample suspected of containing said analyte to provide a measure of said activity;
 - (c) measuring said activity; and
- (d) relating said activity to the presence or absence of said analyte in said sample.
- 37. A method of determining the presence or absence of an analyte of interest in a sample by an homogeneous cloned enzyme

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donor affinity assay comprising:

- enzyme donor fragment attached to a functional surrogate of an analyte of interest, said functional surrogate capable of competing effectively with said analyte for a limiting amount of an affinity receptor for said analyte, said labeled conjugate, on interaction with an enzyme acceptor fragment, exhibiting an activity that is altered in the presence of said affinity receptor and which activity can be measured and related to the amount of said analyte present in a given sample, (ii) said enzyme acceptor fragment, and (iii) said affinity receptor;
- (b) combining said labeled conjugate, enzyme acceptor fragment, and affinity receptor with a sample suspected of containing said analyte to provide a measure of said activity;

(c) measuring said activity; and

- (d) relating said activity to the presence or absence of said analyte in said sample.
 - 38. An affinity assay kit comprising:

(a) a labeled conjugate diposed in a first container means, said labeled conjugate comprising at least one label attached

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to a functional surrogate of an analyte of interest, said functional surrogate capable of competing effectively with said analyte for a limiting amount of an affinity receptor for said analyte, said labeled conjugate capable of exhibiting an activity that is altered on interaction of said labeled conjugate with said affinity receptor and which activity can be measured and related to the amount of said analyte present in a given sample; and

- (b) disposed in a second container means said affinity receptor and, optionally, any substance required for said labeled conjugate to exhibit said activity.
- 39. The kit of claim 38 in which said substance comprises an enzyme substrate.
- 40. The kit of claim 39 in which said substance comprises an enzyme acceptor fragment.
 - 41. The kit of claim 38 in which said activity increases on binding of said labeled conjugate with said affinity receptor.

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42. The kit of claim 38 in which said activity decreases on binding of said labeled conjugate with said affinity receptor.

43. The kit of claim 38 in which said analyte is selected from the group consisting of ferritin, a hepatitis antigen, an antibody against a hepatitis antigen, human chorionic gonadotropin (hCG), human leutinizing hormone (hLH), cytomegalovirus (CMV), chlamydia, streptococcus A, rubella, toxoplasma, herpes, hepatitis, CK-MB, myoglobin, myosin light chain, troponin, carcinoembryonic antigen (CEA), alpha-fetoprotein (AFP), prostrate specific antigen (PSA), CA125 (a tumor marker).

44. A functional surrogate of an analyte of interest comprising a peptide having an immunoreactive group that allows said surrogate to compete effectively with said analyte for a limiting amount of an affinity receptor for said analyte.

45. The functional surrogate of claim 44 in which said peptide comprises about 4 to about 100 amino acid residues.

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46. The functional surrogate of claim 44 which competes effectively with ferritin for a limiting amount of an affinity receptor for ferritin.

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- 47. The functional surrogate of claim 44 which competes effectively with a hepatitis antigen for a limiting amount of an affinity receptor for said hepatitis antigen.
- 48. The functional surrogate of claim 44 in which said antigen is the hepatitis A antigen.
 - 49. The functional surrogate of claim 44 in which said antigen is the hepatitis B antigen.

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- 50. The functional surrogate of claim 44 in which said antigen is the hepatitis C antigen.
- 51. A functional surrogate comprising a peptide having up to about 35 amino acid residues, including the primary sequence motifs depicted in SEQ. ID. NOS. 1-89, 105-115, 127-134, 137-154, 169-180, 193-203, 215-226, 239-247, 255-260, or 267-269.

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- 52. The functional surrogate of claim 51 which further includes 2-10 amino acid residues flanking said sequence motifs.
- 53. The functional surrogate of claim 52 in which said flanking residues are selected among those depicted in said SEQ. ID. NOS.
- 54. A labeled conjugate comprising at least one label attached to a functional surrogate of an analyte of interest, said functional surrogate capable of competing effectively with said analyte for a limiting amount of an affinity receptor for said analyte, said labeled conjugate capable of exhibiting an activity that is altered on interaction of said labeled conjugate with said affinity receptor and which activity can be measured and related to the amount of said analyte present in a given sample.
- 55. A recombinant DNA construct comprising a DNA sequence encoding a functional surrogate of an analyte of interest, said functional surrogate capable of competing effectively with said analyte for a limiting amount of an affinity receptor for said analyte.

56. The construct of claim 53 in which said DNA sequence
is selected from at least those sequences depicted in SEQ. ID. NOS.
90-104, 116-126, 135-136, 155-168, 181-192, 204-214, 227-238,
248-254, 261-266, or 270-272, which encode a primary sequence
motif.

- 57. A transforming vector including the construct of claim 55.
- 58. The vector of claim 57 which is autonomously replicating.
 - 59. Bacteriophage transformed by the vector of claim 57.
- 15 60. A microorganism transformed by the vector of claim 57.
 - 61. A microorganism infected with the bacteriophage of claim 59.

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- 62. A method of obtaining a functional surrogate of an analyte of interest comprising:
- (a) selecting an affinity receptor exhibiting a selective affinity for an analyte of interest;
- (b) screening a random peptide library with said affinity receptor for a binding peptide:
- (c) isolating said binding peptide and identifying its structure.
- 63. The method of claim 62 which further comprises synthesizing said peptide and verifying its capacity to compete with said analyte for a limiting amount of said affinity receptor.
 - 64. The method of claim 62 which further comprises conjugating said peptide to at least one label.
 - 65. The method of claim 62 in which said library is a phage display random peptide library.

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A 63	ACCIPICATIONS AND CHRAINSON ACCITATION		
A. CL.	ASSIFICATION OF SUBJECT MATTER :Please See Extra Sheet.		
1	:Please See Extra Sheet		
	to International Patent Classification (IPC) or to be	oth national classification and IPC	
B. FIE	LDS SEARCHED		
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C. DOC	CUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where	appropriate, of the relevant passages	Relevant to claim No.
X	J.P. RANSOM, "Practical co	ompetitive binding assay	1-5, 8-21, 23-
	methods" published 1976 by T	he C.V. Moshy Company	25, 29-33, and
	(Saint Louis) page 2, see entire of	locument	35
Y	, , , , , , , , , , , , , , , , , , , ,		
			6, 7, 22, 26-28,
		\	34, 36-52, and
			54-56
			34-36
×	CLINICAL IMMUNOLOGY AN	D IMMINOPATHOLOGY	6, 7, 55-65
	Volumne 75, Number 1, issued	April 1995 Dybwad et al	0, 7, 55-65
1	"Structural Characterization of Po	entides That Bind Synovial	44-52 and 54
	Fluid Antibodies From RA Patie	nts: A Novel Strategy for	74-32 and 34
	Identification of Disease-Related	Epitopes Using a Bandom	
ļ	Peptide Library", pages 45-50, se	e entire document	
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X Furthe	er documents are listed in the continuation of Box	C. See patent family annex.	
Spe	cial categories of cited documents:	T later document published after the inter	national filing date or process
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rm PCT/IS.	A/210 (second sheet)(July 1992)*		

International application No. PCT/US96/10498

Cawgory*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
	GENE, Volume 146, issued 1994, Motti et al, "Recognition by Human Sera and Immunogenicity of HBsAg Mimotopes Selected From an M13 Phage Display Library (Hepatitis B virus surface	6, 7, 44, 49, 55-65
	antigen; affinity selection; immuno-screening; serum antibodies; immunization; vaccines; diagnostics) ", pages 191-198, see entire document.	45-52 and 54
	•	

Form PCT/ISA/210 (continuation of second sheet)(July 1992)*

International application No. PCT/US96/10498

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely
2. X Claims Nos.: 53 because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically: because the sequence numbers were not put in the claim after "SEQ. ID. NOS."
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
 As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. X As only some of the required additional search fees were timely paid by the applicant, this international search report cover only those claims for which fees were paid, specifically claims Nos.: 1-43, 44-52,54 (first 10 amino acid sequences), 55-61 (first 10 DNA sequences); 62-65
No required additional search fees were timely paid by the applicant. Consequently, this international search report i restricted to the invention first mentioned in the claims; it is covered by claims Nos.
Remark on Protest The additional search fees were accompanied by the applicant's protest No protest accompanied the payment of additional search fees

Form PCT/[SA/210 (continuation of first sheet(1))(July 1992)*

International application No. PCT/US96/10498

A. CLASSIFICATION OF SUBJECT MATTER: IPC (6):

G01N 33/53, 33/573, 33/537; C07K 7/00, 14/00; C12N 15/09; C12Q 1/70, 1/32

A. CLASSIFICATION OF SUBJECT MATTER: US CL. :

435/5, 7.1, 7.2, 7.4, 7.9, 7.91, 7.92, 26, 320.1, 974, 975; 436/813, 814, 818; 530/324, 325, 326, 327, 328, 329, 330, 350; 935/12, 23, 57, 58, 70, 72

B. FIELDS SEARCHED

Minimum documentation searched Classification System: U.S.

435/5, 7.1, 7.2, 7.4, 7.9, 7.91, 7.92, 26, 320.1, 974, 975; 436/813, 814, 818; 530/324, 325, 326, 327, 328, 329, 330, 350; 935/12, 23, 57, 58, 70, 72